



- (51) International Patent Classification:
C07K 14/495 (2006.01) *C12N 15/62* (2006.01)
- (21) International Application Number:
PCT/US2014/049254
- (22) International Filing Date:
31 July 2014 (31.07.2014)
- (25) Filing Language: English
- (26) Publication Language: English
- (30) Priority Data:
61/860,723 31 July 2013 (31.07.2013) US
- (71) Applicant: **AMGEN INC.** [US/US]; One Amgen Center Dr., Thousand Oaks, CA 91320 (US).
- (72) Inventors: **XIONG, Yumei**; 140 Daley Court, San Bruno, CA 94066 (US). **ZHANG, Yi**; 541 Driscoll Place, Palo Alto, CA 94306 (US). **SHENG, Jackie, Z.**; 559 Lone Oak Drive, Thousand Oaks, CA 91362 (US). **HAMBURGER, Agnes, Eva**; 3885 Calle Alta Vista, Newbury Park, CA 91320 (US). **VENIANT-ELLISON, Murielle**; 645 Camino Manzanitas, Thousand Oaks, CA 91360 (US). **SHIM-AMOTO, Grant**; 368 Via Colinas, Westlake Village, CA 91362 (US). **MIN, Xiaoshan**; 2108 Clarice Lane, Burlingame, CA 94010 (US). **WANG, Zhulun**; 660 Glenbrook Drive, Palo Alto, CA 94306 (US). **TANG, Jie**; 922 Moreno Avenue, Palo Alto, CA 94303 (US). **KANNAN, Gunasekaran**; 1056 Amberton Lane, Newbury Park, CA 91320 (US). **MOCK, Marissa**; 229 S Dewey Avenue, Newbury Park, CA 91320 (US). **WALKER, Kenneth**; 175 Mesa Avenue, Newbury Park, CA 91320 (US).

- (74) Agent: **LESCHENSKY, William**; 1120 Veterans Blvd, San Francisco, CA 94080 (US).

- (81) Designated States (unless otherwise indicated, for every kind of national protection available): AE, AG, AL, AM, AO, AT, AU, AZ, BA, BB, BG, BH, BN, BR, BW, BY, BZ, CA, CH, CL, CN, CO, CR, CU, CZ, DE, DK, DM, DO, DZ, EC, EE, EG, ES, FI, GB, GD, GE, GH, GM, GT, HN, HR, HU, ID, IL, IN, IR, IS, JP, KE, KG, KN, KP, KR, KZ, LA, LC, LK, LR, LS, LT, LU, LY, MA, MD, ME, MG, MK, MN, MW, MX, MY, MZ, NA, NG, NI, NO, NZ, OM, PA, PE, PG, PH, PL, PT, QA, RO, RS, RU, RW, SA, SC, SD, SE, SG, SK, SL, SM, ST, SV, SY, TH, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, ZA, ZM, ZW.

- (84) Designated States (unless otherwise indicated, for every kind of regional protection available): ARIPO (BW, GH, GM, KE, LR, LS, MW, MZ, NA, RW, SD, SL, SZ, TZ, UG, ZM, ZW), Eurasian (AM, AZ, BY, KG, KZ, RU, TJ, TM), European (AL, AT, BE, BG, CH, CY, CZ, DE, DK, EE, ES, FI, FR, GB, GR, HR, HU, IE, IS, IT, LT, LU, LV, MC, MK, MT, NL, NO, PL, PT, RO, RS, SE, SI, SK, SM, TR), OAPI (BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, KM, ML, MR, NE, SN, TD, TG).

Published:

- with international search report (Art. 21(3))
- before the expiration of the time limit for amending the claims and to be republished in the event of receipt of amendments (Rule 48.2(h))
- with sequence listing part of description (Rule 5.2(a))

- (54) Title: GROWTH DIFFERENTIATION FACTOR 15 (GDF-15) CONSTRUCTS

(57) Abstract: Constructs comprising GDF15, and mutants thereof are provided. In various embodiments the constructs comprising GDF15, and mutants thereof, can be of use in the treatment or ameliorating a metabolic disorder. In various embodiments the metabolic disease or disorder is type 2 diabetes, obesity, dyslipidemia, elevated glucose levels, elevated insulin levels and diabetic nephropathy.



GROWTH DIFFERENTIATION FACTOR 15 (GDF-15) CONSTRUCTS

SEQUENCE LISTING

The instant application contains a Sequence Listing which has been submitted electronically in ASCII format and is hereby incorporated by reference in its entirety. Said ASCII copy, created on July 29, 2014, is named A-1850-WO-PCT_SL.txt and is 630,200 bytes in size.

FIELD OF THE INVENTION

The instant disclosure relates to monomers and multimers comprising a polypeptide comprising a GDF15 region.

BACKGROUND OF THE INVENTION

Growth differentiation factor 15 (GDF15) is a divergent member of the TGF β superfamily. It is also called macrophage inhibitory cytokine 1 (MIC1) (Bootcov MR, 1997, *Proc Natl Acad Sci* 94:11514-9), placental bone morphogenetic factor (PLAB) (Hromas R 1997, *Biochim Biophys Acta*. 1354:40-4), placental transforming growth factor beta (PTGFB) (Lawton LN 1997, *Gene*. 203:17-26), prostate derived factor (PDF) (Paralkar VM 1998, *J Biol Chem*. 273:13760-7), and nonsteroidal anti-inflammatory drug-activated gene (NAG-1) (Baek SJ 2001, *J Biol Chem*. 276: 33384-92).

Human GDF15 gene is located on chromosome 19p13.2-13.1; rat GDF15 gene is located on chromosome 16; and mouse GDF15 gene is located on chromosome 8. The GDF15 open reading frames span two exons (Bottner M 1999, *Gene*. 237:105-11 and NCBI). The mature GDF15 peptide shares low homology with other family members (Katoh M 2006, *Int J Mol Med*. 17:951-5.).

GDF15 is synthesized as a large precursor protein that is cleaved at the dibasic cleavage site to release the carboxyterminal mature peptide. The mouse and rat GDF15 prepro-peptides both contain 303 amino acids. Human full-length precursor contains 308 amino acids. The rodent mature peptides contain 115 amino acids after processing at the RGRR (SEQ ID NO:1) cleavage site. The human mature peptide contains 112 amino acids after processing at the RGRRRAR (SEQ ID NO:2) cleavage site. Human mature GDF15 peptide shares 66.1% and 68.1% sequence similarity with rat and mouse mature GDF15 peptides (Bottner M 1999, *Gene*. 237:105-11; Bauskin AR 2000, *EMBO J*. 19:2212-20; NCBI). There is no glycosylation site in the mature GDF15 peptide.

The mature GDF15 peptide contains the seven conserved cysteine residues required for the formation of the cysteine knot motif (having three intrachain disulfide bonds) and the single interchain disulfide bond that are typical for TGF β superfamily members. The mature

GDF15 peptide further contains two additional cysteine residues that form a fourth intrachain disulfide bond. Biologically active GDF15 is a 25KD homodimer of the mature peptide covalently linked by one interchain disulfide bond.

GDF15 circulating levels have been reported to be elevated in multiple pathological and physiological conditions, most notably pregnancy (Moore AG 2000, *J Clin Endocrinol Metab* 85: 4781-4788), β -thalassemia (Tanno T 2007, *Nat Med* 13:1096-101; Zimmermann MB, 2008 *Am J Clin Nutr* 88:1026-31), and congenital dyserythropoietic anemia (Tamary H 2008, *Blood*. 112:5241-4). GDF15 has also been linked to multiple biological activities in literature reports. Studies of GDF15 knockout and transgenic mice suggested that GDF15 may be protective against ischemic/reperfusion- or overload-induced heart injury (Kempf T, 2006, *Circ Res*.98:351-60; Xu J, 2006, *Circ Res*. 98:342-50), protective against aging-associated motor neuron and sensory neuron loss (Strelau J, 2009, *J Neurosci*. 29 :13640-8), mildly protective against metabolic acidosis in kidney, and may cause cachexia in cancer patients (Johnen H 2007 *Nat Med*. 11:1333-40). Many groups also have studied the role of GDF15 in cell apoptosis and proliferation and reported controversial results using different cell culture and xenograft models. Studies on transgenic mice showed that GDF15 is protective against carcinogen or Apc mutation induced neoplasia in intestine and lung (Baek SJ 2006, *Gastroenterology*. 131:1553-60; Cekanova M 2009, *Cancer Prev Res* 2:450-8).

SUMMARY OF THE INVENTION

Provided herein are fusion proteins comprising a GDF15 polypeptide or a GDF15 mutant polypeptide and an Fc domain.

In a one embodiment, the Fc domain comprises a sequence selected from the group consisting of SEQ ID NOs:16, 22, 28, 29, 33, 35, 38, 48, 85, 91, 106, 132, 141, 148, 155, 162, 169, 176, 183, 192, 199, 206, 213, 220, 227, 233, 236, 268, 275, 281, 282, 283, 284, 285, 286, 287, 288, 289, 290, 291, 292, 293, 294, 295, 296, 297, 298, 299, 300, 301 and 302. In another embodiment, the GDF15 polypeptide or a GDF15 mutant polypeptide comprises a sequence selected from the group consisting of SEQ ID NOs:4, 8, 12, 25, 52 and 55. In another embodiment, the fusion protein further comprises a polypeptide linker. In one embodiment, the polypeptide linker has a sequence selected from the group consisting of SEQ ID NOs:18, 30, 34, 40, 58, 61, 64, 69, 72, 75, 78, 113, 116, 119, 122, 125, 128. In one embodiment, the fusion protein comprises two or more Fc domains. In one embodiment, the fusion protein comprises two or more polypeptide linkers. In one embodiment, the fusion protein comprises a sequence selected from the group consisting of SEQ ID NOs: 46, 24, 27, 32, 37, 20, 42, 50, 54, 57, 60, 63, 66, 68, 71, 74, 77, 82, 84, 88, 93, 96, 98, 100, 102, 104, 108, 134, 137, 139, 143, 146, 150, 153, 269, 272, 276, 279, 157, 160, 164, 167, 171, 174, 178, 181, 185, 188, 194, 197, 201, 204, 208, 211, 215, 218, 222, 225, 229, 232, 233, 238 and 240.

Also provided herein are dimers comprising (i) a first polypeptide chain comprising one of the foregoing fusion proteins, and (ii) a second polypeptide chain comprising an Fc domain. In yet a further embodiment, the construct further comprises a sequence selected from the group consisting of SEQ ID NOs: 16, 22, 28, 29, 33, 35, 38, 48, 85, 91, 106, 132,
5 141, 148, 155, 162, 169, 176, 183, 192, 199, 206, 213, 220, 227, 233, 236, 268, 275, 281, 282, 283, 284, 285, 286, 287, 288, 289, 290, 291, 292, 293, 294, 295, 296, 297, 298, 299, 300, 301 and 302.

In one embodiment, the first and second polypeptide chains are non-covalently associated. In another embodiment, the first and second polypeptide chains are covalently
10 associated. In one embodiment, the first and second polypeptide chains are covalently associated via disulfide bonds between their respective Fc domains. In another embodiment, the first and second polypeptide chains are associated by both covalent and non-covalent interactions.

In a particular embodiment, a dimer is provided comprising: (a) two fusion proteins
15 comprising the sequence of SEQ ID NO:46; (b) two fusion proteins comprising the sequence of SEQ ID NO:24; or (c) two fusion proteins comprising the sequence of SEQ ID NO:27

In a particular embodiment, a dimer is provided comprising (a) two fusion proteins comprising the sequence of SEQ ID NO:32; or (b) two fusion proteins comprising the sequence of SEQ ID NO:37;

20 In a particular embodiment, a dimer is provided comprising a first polypeptide chain comprising the sequence of SEQ ID NO:20 and a second polypeptide chain comprising the sequence of SEQ ID NO:17.

In a particular embodiment, a dimer is provided comprising (a) a first polypeptide chain comprising the sequence of SEQ ID NO:42 and a second polypeptide chain comprising
25 the sequence of SEQ ID NO:39; (b) a first polypeptide chain comprising the sequence of SEQ ID NO:50 and a second polypeptide chain comprising the sequence of SEQ ID NO:47; (c) a first polypeptide chain comprising the sequence of SEQ ID NO:54 and a second polypeptide chain comprising the sequence of SEQ ID NO:47; (d) a first polypeptide chain comprising the sequence of SEQ ID NO:57 and a second polypeptide chain comprising the sequence of SEQ
30 ID NO:47; (e) a first polypeptide chain comprising the sequence of SEQ ID NO:60 and a second polypeptide chain comprising the sequence of SEQ ID NO:47; (f) a first polypeptide chain comprising the sequence of SEQ ID NO:63 and a second polypeptide chain comprising the sequence of SEQ ID NO:47; (g) a first polypeptide chain comprising the sequence of SEQ ID NO:66 and a second polypeptide chain comprising the sequence of SEQ ID NO:47; (h) a
35 first polypeptide chain comprising the sequence of SEQ ID NO:68 and a second polypeptide chain comprising the sequence of SEQ ID NO:47; (i) a first polypeptide chain comprising the sequence of SEQ ID NO:71 and a second polypeptide chain comprising the sequence of SEQ

ID NO:47; (j) a first polypeptide chain comprising the sequence of SEQ ID NO:74 and a second polypeptide chain comprising the sequence of SEQ ID NO:47; (k) a first polypeptide chain comprising the sequence of SEQ ID NO:77 and a second polypeptide chain comprising the sequence of SEQ ID NO:47; (l) a first polypeptide chain comprising the sequence of SEQ ID NO:80 and a second polypeptide chain comprising the sequence of SEQ ID NO:47; (m) a first polypeptide chain comprising the sequence of SEQ ID NO:82 and a second polypeptide chain comprising the sequence of SEQ ID NO:47; or (n) a first polypeptide chain comprising the sequence of SEQ ID NO:84 and a second polypeptide chain comprising the sequence of SEQ ID NO:47.

10 In a particular embodiment, a dimer is provided comprising (a) a first polypeptide chain comprising the sequence of SEQ ID NO:88 and a second polypeptide chain comprising the sequence of SEQ ID NO:86; (b) a first polypeptide chain comprising the sequence of SEQ ID NO:93 and a second polypeptide chain comprising the sequence of SEQ ID NO:90; (c) a first polypeptide chain comprising the sequence of SEQ ID NO:96 and a second polypeptide chain comprising the sequence of SEQ ID NO:90; (d) a first polypeptide chain comprising the sequence of SEQ ID NO:98 and a second polypeptide chain comprising the sequence of SEQ ID NO:90; (e) a first polypeptide chain comprising the sequence of SEQ ID NO:100 and a second polypeptide chain comprising the sequence of SEQ ID NO:90; (f) a first polypeptide chain comprising the sequence of SEQ ID NO:102 and a second polypeptide chain comprising the sequence of SEQ ID NO:90; (g) a first polypeptide chain comprising the sequence of SEQ ID NO:104 and a second polypeptide chain comprising the sequence of SEQ ID NO:90; or (h) a first polypeptide chain comprising the sequence of SEQ ID NO:108 and a second polypeptide chain comprising the sequence of SEQ ID NO:105.

25 In a particular embodiment, a dimer is provided comprising (a) a first polypeptide chain comprising the sequence of SEQ ID NO:112 and a second polypeptide chain comprising the sequence of SEQ ID NO:12; (b) two polypeptide chains, each comprising the sequence of SEQ ID NO:112; (c) two polypeptide chains, each comprising the sequence of SEQ ID NO:115; (d) two polypeptide chains, each comprising the sequence of SEQ ID NO:118; (e) two polypeptide chains, each comprising the sequence of SEQ ID NO:121; (f) two polypeptide chains, each comprising the sequence of SEQ ID NO:124; (g) two polypeptide chains, each comprising the sequence of SEQ ID NO:127; (h) two polypeptide chains, each comprising the sequence of SEQ ID NO:130; or (i) two polypeptide chains, each comprising the sequence of SEQ ID NO:242.

35 In a particular embodiment, a dimer is provided comprising: (a) a first polypeptide chain comprising the sequence of SEQ ID NO:134 and a second polypeptide chain comprising the sequence of SEQ ID NO:131; (b) a first polypeptide chain comprising the sequence of SEQ ID NO:137 and a second polypeptide chain comprising the sequence of

SEQ ID NO:131; (c) a first polypeptide chain comprising the sequence of SEQ ID NO:139 and a second polypeptide chain comprising the sequence of SEQ ID NO:131; (d) a first polypeptide chain comprising the sequence of SEQ ID NO:143 and a second polypeptide chain comprising the sequence of SEQ ID NO:140; (e) a first polypeptide chain comprising the sequence of SEQ ID NO:146 and a second polypeptide chain comprising the sequence of SEQ ID NO:140; (f) a first polypeptide chain comprising the sequence of SEQ ID NO:150 and a second polypeptide chain comprising the sequence of SEQ ID NO:147; (g) a first polypeptide chain comprising the sequence of SEQ ID NO:153 and a second polypeptide chain comprising the sequence of SEQ ID NO:147; (h) a first polypeptide chain comprising the sequence of SEQ ID NO:269 and a second polypeptide chain comprising the sequence of SEQ ID NO:267; (i) a first polypeptide chain comprising the sequence of SEQ ID NO:272 and a second polypeptide chain comprising the sequence of SEQ ID NO:267; (j) a first polypeptide chain comprising the sequence of SEQ ID NO:276 and a second polypeptide chain comprising the sequence of SEQ ID NO:274; or (k) a first polypeptide chain comprising the sequence of SEQ ID NO:279 and a second polypeptide chain comprising the sequence of SEQ ID NO:274.

In a particular embodiment, a dimer is provided comprising: (a) a first polypeptide chain comprising the sequence of SEQ ID NO:157 and a second polypeptide chain comprising the sequence of SEQ ID NO:154; (b) a first polypeptide chain comprising the sequence of SEQ ID NO:160 and a second polypeptide chain comprising the sequence of SEQ ID NO:154; (c) a first polypeptide chain comprising the sequence of SEQ ID NO:164 and a second polypeptide chain comprising the sequence of SEQ ID NO:161; (d) a first polypeptide chain comprising the sequence of SEQ ID NO:167 and a second polypeptide chain comprising the sequence of SEQ ID NO:161.

In a particular embodiment, a dimer is provided comprising: (a) a first polypeptide chain comprising the sequence of SEQ ID NO:171 and a second polypeptide chain comprising the sequence of SEQ ID NO:168; or (b) a first polypeptide chain comprising the sequence of SEQ ID NO:174 and a second polypeptide chain comprising the sequence of SEQ ID NO:168

In a particular embodiment, a dimer is provided comprising: (a) a first polypeptide chain comprising the sequence of SEQ ID NO:178 and a second polypeptide chain comprising the sequence of SEQ ID NO:175; (b) a first polypeptide chain comprising the sequence of SEQ ID NO:181 and a second polypeptide chain comprising the sequence of SEQ ID NO:175; (c) a first polypeptide chain comprising the sequence of SEQ ID NO:185 and a second polypeptide chain comprising the sequence of SEQ ID NO:182; (d) a first polypeptide chain comprising the sequence of SEQ ID NO:188 and a second polypeptide chain comprising the sequence of SEQ ID NO:182; (e) a first polypeptide chain comprising

the sequence of SEQ ID NO:194 and a second polypeptide chain comprising the sequence of SEQ ID NO:191; (f) a first polypeptide chain comprising the sequence of SEQ ID NO:197 and a second polypeptide chain comprising the sequence of SEQ ID NO:191; (g) a first polypeptide chain comprising the sequence of SEQ ID NO:201 and a second polypeptide chain comprising the sequence of SEQ ID NO:198; or (h) a first polypeptide chain comprising the sequence of SEQ ID NO:204 and a second polypeptide chain comprising the sequence of SEQ ID NO:198.

In a particular embodiment, a dimer is provided comprising: (a) a first polypeptide chain comprising the sequence of SEQ ID NO:208 and a second polypeptide chain comprising the sequence of SEQ ID NO:205; (b) a first polypeptide chain comprising the sequence of SEQ ID NO:211 and a second polypeptide chain comprising the sequence of SEQ ID NO:205; (c) a first polypeptide chain comprising the sequence of SEQ ID NO:215 and a second polypeptide chain comprising the sequence of SEQ ID NO:212; or (d) a first polypeptide chain comprising the sequence of SEQ ID NO:218 and a second polypeptide chain comprising the sequence of SEQ ID NO:212.

In a particular embodiment, a dimer is provided comprising: (a) a first polypeptide chain comprising the sequence of SEQ ID NO:222 and a second polypeptide chain comprising the sequence of SEQ ID NO:219; (b) a first polypeptide chain comprising the sequence of SEQ ID NO:225 and a second polypeptide chain comprising the sequence of SEQ ID NO:219; (c) a first polypeptide chain comprising the sequence of SEQ ID NO:229 and a second polypeptide chain comprising the sequence of SEQ ID NO:226; or (d) a first polypeptide chain comprising the sequence of SEQ ID NO:232 and a second polypeptide chain comprising the sequence of SEQ ID NO:226.

In a particular embodiment, a dimer is provided comprising: (a) two polypeptide chains, each comprising the sequence of SEQ ID NO:235; (b) two polypeptide chains, each comprising the sequence of SEQ ID NO:238; or (c) two polypeptide chains, each comprising the sequence of SEQ ID NO:240.

In certain embodiments, a tetramer is provided, comprising (i) a first dimer comprising one of the foregoing dimers and (ii) a second dimer comprising one of the foregoing dimers. In certain embodiments, the first polypeptide chain of the first dimer is linked to the first polypeptide chain of the second dimer via an interchain disulfide bond between their respective GDF15 regions.

In some embodiments, the dimer is not selected from the group comprising DhCpmFc(-)-(G4S)₄-GDF15:DhCpmFc(+), DhCpmFc(+)-(G4S)₄-GDF15:DhCpmFc(-), DhCpmFc(-)-(G4S)₄-GDF15(H6D):DhCpmFc(+), DhCpmFc(+)-(G4S)₄-GDF15(H6D):DhCpmFc(-), DhCpmFc(+)-(G4S)₄-GDF15(N3Q):DhCpmFc(-), DhCpmFc(+)-GDF15:DhCpmFc(-), DhCpmFc(+)-G4-GDF15:DhCpmFc(-), DhCpmFc(+)-(G4S)₂-

GDF15:DhCpmFc(-), DhCpmFc(+)-(G₄Q)₄-GDF15:DhCpmFc(-), DhCpmFc(+)(L351C)-G₄-GDF15:DhCpmFc(-)(L351C), DhCpmFc(+)(S354C)-G₄-GDF15:DhCpmFc(-)(Y349C), CpmFc(-)-(G₄S)₄-GDF15:CpmFc(+), Fc-(G₄S)₈-Fc-GS(G₄S)₄-GDF15, Fc-(G₄S)₃-Fc-GS(G₄S)₄-GDF15 and Fc-(G₄S)₅-Fc-GS(G₄S)₄-GDF15.

- 5 In some embodiments, the dimer is not selected from the group consisting of DhCpmFc(-)-(G₄S)₄-GDF15:DhCpmFc(+), DhCpmFc(+)-(G₄S)₄-GDF15:DhCpmFc(-), DhCpmFc(-)-(G₄S)₄-GDF15(H6D):DhCpmFc(+), DhCpmFc(+)-(G₄S)₄-GDF15(H6D):DhCpmFc(-), DhCpmFc(+)-(G₄S)₄-GDF15(N3Q):DhCpmFc(-), DhCpmFc(+)-GDF15:DhCpmFc(-), DhCpmFc(+)-G₄-GDF15:DhCpmFc(-), DhCpmFc(+)-(G₄S)₂-GDF15:DhCpmFc(-), DhCpmFc(+)-(G₄Q)₄-GDF15:DhCpmFc(-), DhCpmFc(+)(L351C)-G₄-GDF15:DhCpmFc(-)(L351C) and DhCpmFc(+)(S354C)-G₄-GDF15:DhCpmFc(-)(Y349C).

In some embodiments, the fusion protein is not a selected from the group consisting of Fc-(G₄S)₈-Fc-GS(G₄S)₄-GDF15, Fc-(G₄S)₃-Fc-GS(G₄S)₄-GDF15 and Fc-(G₄S)₅-Fc-GS(G₄S)₄-GDF15.

- 15 In some embodiments, the dimer is not selected from the group consisting of DhCpmFc(+)(L351C)-G₄-GDF15:DhCpmFc(-)(L351C) and DhCpmFc(+)(S354C)-G₄-GDF15:DhCpmFc(-)(Y349C).

- In some embodiments, the dimer is not DhCpmFc(-)-(G₄S)₄-GDF15:DhCpmFc(+), DhCpmFc(+)-(G₄S)₄-GDF15:DhCpmFc(-), DhCpmFc(-)-(G₄S)₄-GDF15(H6D):DhCpmFc(+), DhCpmFc(+)-(G₄S)₄-GDF15(H6D):DhCpmFc(-), DhCpmFc(+)-(G₄S)₄-GDF15(N3Q):DhCpmFc(-), DhCpmFc(+)-GDF15:DhCpmFc(-), DhCpmFc(+)-G₄-GDF15:DhCpmFc(-), DhCpmFc(+)-(G₄S)₂-GDF15:DhCpmFc(-), DhCpmFc(+)-(G₄Q)₄-GDF15:DhCpmFc(-), DhCpmFc(+)(L351C)-G₄-GDF15:DhCpmFc(-)(L351C), DhCpmFc(+)(S354C)-G₄-GDF15:DhCpmFc(-)(Y349C), CpmFc(-)-(G₄S)₄-GDF15:CpmFc(+), Fc-(G₄S)₈-Fc-GS(G₄S)₄-GDF15, Fc-(G₄S)₃-Fc-GS(G₄S)₄-GDF15 or Fc-(G₄S)₅-Fc-GS(G₄S)₄-GDF15.

- In some embodiments, the dimer is not DhCpmFc(-)-(G₄S)₄-GDF15:DhCpmFc(+), DhCpmFc(+)-(G₄S)₄-GDF15:DhCpmFc(-), DhCpmFc(-)-(G₄S)₄-GDF15(H6D):DhCpmFc(+), DhCpmFc(+)-(G₄S)₄-GDF15(H6D):DhCpmFc(-), DhCpmFc(+)-(G₄S)₄-GDF15(N3Q):DhCpmFc(-), DhCpmFc(+)-GDF15:DhCpmFc(-), DhCpmFc(+)-G₄-GDF15:DhCpmFc(-), DhCpmFc(+)-(G₄S)₂-GDF15:DhCpmFc(-), DhCpmFc(+)-(G₄Q)₄-GDF15:DhCpmFc(-), DhCpmFc(+)(L351C)-G₄-GDF15:DhCpmFc(-)(L351C) or DhCpmFc(+)(S354C)-G₄-GDF15:DhCpmFc(-)(Y349C).

- In some embodiments, the fusion protein is not selected from the group consisting of Fc-(G₄S)₈-Fc-GS(G₄S)₄-GDF15, Fc-(G₄S)₃-Fc-GS(G₄S)₄-GDF15 and Fc-(G₄S)₅-Fc-GS(G₄S)₄-GDF15.

In some embodiments, the fusion protein is not selected from the group consisting of DhCpmFc(+)(L351C)-G₄-GDF15:DhCpmFc(-)(L351C) and DhCpmFc(+)(S354C)-G₄-GDF15:DhCpmFc(-)(Y349C).

In some embodiments, the dimer is not DhCpmFc(-)-(G₄S)₄-GDF15:DhCpmFc(+). In some
 5 some embodiments, the dimer is not DhCpmFc(+)-(G₄S)₄-GDF15:DhCpmFc(-). In some
 embodiments, the dimer is not DhCpmFc(-)-(G₄S)₄-GDF15(H6D):DhCpmFc(+). In some
 embodiments, the dimer is not DhCpmFc(+)-(G₄S)₄-GDF15(H6D):DhCpmFc(-). In some
 embodiments, the dimer is not DhCpmFc(+)-(G₄S)₄-GDF15(N3Q):DhCpmFc(-). In some
 embodiments, the dimer is not DhCpmFc(+)-GDF15:DhCpmFc(-). In some embodiments, the
 10 dimer is not DhCpmFc(+)-G₄-GDF15:DhCpmFc(-). In some embodiments, the dimer is not
 DhCpmFc(+)-(G₄S)₂-GDF15:DhCpmFc(-). In some embodiments, the dimer is not
 DhCpmFc(+)-(G₄Q)₄-GDF15:DhCpmFc(-). In some embodiments, the dimer is not
 DhCpmFc(+)(L351C)-G₄-GDF15:DhCpmFc(-)(L351C). In some embodiments, the dimer is
 not DhCpmFc(+)(S354C)-G₄-GDF15:DhCpmFc(-)(Y349C). In some embodiments, the dimer
 15 is not CpmFc(-)-(G₄S)₄-GDF15:CpmFc(+). In some embodiments, the dimer is not Fc-(G₄S)₈-
 Fc-GS(G₄S)₄-GDF15. In some embodiments, the dimer is not Fc-(G₄S)₃-Fc-GS(G₄S)₄-
 GDF15. In some embodiments, the dimer is not Fc-(G₄S)₅-Fc-GS(G₄S)₄-GDF15.

Also provided herein are fusion proteins comprising a GDF15 polypeptide or a
 GDF15 mutant polypeptide and a human serum albumin (HSA) polypeptide. In a further
 20 embodiment, the HSA polypeptide the sequence of SEQ ID NO:110. In another embodiment,
 the GDF15 polypeptide or a GDF15 mutant polypeptide comprises a sequence selected from
 the group consisting of SEQ ID NOs:4, 8, 12, 25, 52 and 55. In another embodiment, the
 fusion protein further comprises a polypeptide linker, joining the GDF15 polypeptide or
 GDF15 mutant polypeptide to the HSA polypeptide. In one embodiment, the polypeptide
 25 linker has a sequence selected from the group consisting of SEQ ID NOs:18, 30, 34, 40, 58,
 61, 64, 69, 72, 75, 78, 113, 116, 119, 122, 125, 128. In one embodiment, the fusion protein
 comprises two or more HSA polypeptides. In one embodiment, the fusion protein comprises a
 sequence selected from the group consisting of SEQ ID NOs: 115, 118, 121, 124, 127 and
 130.

Also provided herein are dimers comprising (i) a first polypeptide chain comprising a
 fusion protein comprising a first GDF15 region and a first HSA polypeptide, and (ii) a second
 polypeptide chain comprising a second HSA polypeptide. In some embodiments, the second
 polypeptide chain further comprises a second GDF15 region. In some embodiments, the
 dimer is a heterodimer (i.e., the first and second polypeptide chain have different sequences).
 35 In some embodiments, the dimer is a homodimer (i.e., the first and second polypeptide chain
 have the same sequence). In some embodiments, the first polypeptide chain comprises a
 sequence selected from the group consisting of SEQ ID NOs: 115, 118, 121, 124, 127 and

130. In some embodiments, the second polypeptide chain comprises a sequence selected from the group consisting of SEQ ID NOs: 115, 118, 121, 124, 127 and 130. In one embodiment, the first and second polypeptide chains are non-covalently associated. In another embodiment, the first and second polypeptide chains are covalently associated. In one embodiment, the second polypeptide chain comprises a second GDF15 region and first and second polypeptide chains are covalently associated via disulfide bonds between their respective GDF15 regions. In another embodiment, the first and second polypeptide chains are associated by both covalent and non-covalent interactions.

Also provided herein are dimers comprising (i) a first polypeptide chain comprising a fusion protein comprising a first GDF15 region and an HSA polypeptide, and (ii) a second polypeptide chain comprising a second GDF15 region. In some embodiments, the first polypeptide chain comprises a sequence selected from the group consisting of SEQ ID NOs: 115, 118, 121, 124, 127 and 130. In some embodiments, the second polypeptide chain comprises a sequence selected from the group consisting of SEQ ID NOs: 4, 8, 12, 25, 52 and 55. In one embodiment, the first and second polypeptide chains are non-covalently associated. In another embodiment, the first and second polypeptide chains are covalently associated. In another embodiment, the first and second polypeptide chains are associated by both covalent and non-covalent interactions.

BRIEF DESCRIPTION OF THE DRAWINGS

Figure 1 is a graphic depicting of a knob-hole construct comprising a dimer of two DhknobFc-(G₄S)₄-GDF15:DhholeFc heterodimers.

Figure 2 is a graphic depicting a DhMonoFc construct comprising a dimer of two DhMonoFc-(G₄S)₄-GDF15 fusion proteins.

Figure 3 is a graphic depicting a HemiFc construct comprising a dimer of two GGGFc-(G₄S)₄-Fc-S(G₄S)₄-GDF15 fusion proteins.

Figure 4 is a graphic depicting a charged pair (delHinge) construct comprising a dimer of two DhCpmFc(-)-(G₄S)₂-GDF15:DhCpmFc(+) heterodimers.

Figure 5 is a graphic depicting a charged pair (delHinge) cysteine clamp construct comprising a dimer of two DhCpmFc(-)(L351C)-(G₄S)₂-GDF15:DhCpmFc(-)(L351C) heterodimers.

Figure 6 is a graphic depicting an HSA construct comprising a dimer of two HSA-(G₄S)₄-GDF15 fusion proteins.

Figure 7 is a plot showing the effect on food intake (g food/g body weight (BW)) of ob/ob mice as a function of dose (log [g construct/kg BW]) using a dimer of the DhknobFc-(G₄S)₄-GDF15:DhholeFc heterodimer.

Figure 8 is a plot showing the effect on food intake (g food/g body weight) of ob/ob mice as a function of dose (log [g construct/kg BW]) using a dimer of the Fc-(G₄S)₄-GDF15 fusion protein.

Figure 9 is a plot showing the effect on food intake (g food/g body weight) of ob/ob mice as a function of dose (log [g construct/kg BW]) using a dimer of the Fc-(G₄S)₄-GDF15(H6D) fusion protein.

Figure 10 is a plot showing the effect on food intake (g food/g body weight) of ob/ob mice as a function of dose (log [g construct/kg BW]) using a dimer of the Fc-(G₄S)₄-Fc-(G₄S)₄-GDF15 fusion protein.

Figure 11 is a plot showing the effect on food intake (g food/g body weight) of ob/ob mice as a function of dose (log [g construct/kg BW]) using a dimer of the DhFc-(G₄S)₅-DhFc-(G₄S)₄-GDF15 fusion protein.

Figure 12 is a plot showing the effect on food intake (g food/g body weight) of ob/ob mice as a function of dose (log [g construct/kg BW]) using a dimer of the DhCpmFc(+)-(1K)-GDF15:DhCpmFc(-) heterodimer.

Figure 13 is a plot showing the effect on food intake (g food/g body weight) of ob/ob mice as a function of dose (log [g construct/kg BW]) using a dimer of the DhCpmFc(-)-GDF15:DhCpmFc(+) heterodimer.

Figure 14 is a plot showing the effect on food intake (g food/g body weight) of ob/ob mice as a function of dose (log [g construct/kg BW]) using a dimer of the DhCpmFc(-)-GDF15(N3D):DhCpmFc(+) heterodimer.

Figure 15 is a plot showing the effect on food intake (g food/g body weight) of ob/ob mice as a function of dose (log [g construct/kg BW]) using a dimer of the DhCpmFc(-)-GDF15(Ndel3):DhCpmFc(+) heterodimer.

Figure 16 is a plot showing the effect on food intake (g food/g body weight) of ob/ob mice as a function of dose (log [g construct/kg BW]) using a dimer of the DhCpmFc(-)-G₄-GDF15(N3D):DhCpmFc(+) heterodimer.

Figure 17 is a plot showing the effect on food intake (g food/g body weight) of ob/ob mice as a function of dose (log [g construct/kg BW]) using a dimer of the DhCpmFc(-)-G₄S-GDF15:DhCpmFc(+) heterodimer.

Figure 18 is a plot showing the effect on food intake (g food/g body weight) of ob/ob mice as a function of dose (log [g construct/kg BW]) using a dimer of the DhCpmFc(-)-(G₄S)₂-GDF15:DhCpmFc(+) heterodimer.

Figure 19 is a plot showing the effect on food intake (g food/g body weight) of ob/ob mice as a function of dose (log [g construct/kg BW]) using a dimer of the DhCpmFc(-)-(G₄S)₂-GDF15(N3D):DhCpmFc(+) heterodimer.

Figure 20 is a plot showing the effect on food intake (g food/g body weight) of ob/ob mice as a function of dose (log [g construct/kg BW]) using a dimer of the DhCpmFc(-)-(G₄Q)₂-GDF15(N3D):DhCpmFc(+) heterodimer.

Figure 21 is a plot showing the effect on food intake (g food/g body weight) of ob/ob mice as a function of dose (log [g construct/kg BW]) using a dimer of the DhCpmFc(-)-G₄P-GDF15:DhCpmFc(+) heterodimer.

Figure 22 is a plot showing the effect on food intake (g food/g body weight) of ob/ob mice as a function of dose (log [g construct/kg BW]) using a dimer of the DhCpmFc(-)-(G₄P)₂-GDF15:DhCpmFc(+) heterodimer.

Figure 23 is a plot showing the effect on food intake (g food/g body weight) of ob/ob mice as a function of dose (log [g construct/kg BW]) using a dimer of the DhCpmFc(-)-G₄Q-GDF15:DhCpmFc(+) heterodimer.

Figure 24 is a plot showing the effect on food intake (g food/g body weight) of ob/ob mice as a function of dose (log [g construct/kg BW]) using a dimer of the DhCpmFc(-)-(G₄Q)₂-GDF15(N3D):DhCpmFc(+) heterodimer.

Figure 25 is a plot showing the effect on food intake (g food/g body weight) of ob/ob mice as a function of dose (log [g construct/kg BW]) using a dimer of the DhCpmFc(-)-(G₄Q)₂-GDF15(Ndel3):DhCpmFc(+) heterodimer.

Figure 26 is a plot showing the effect on food intake (g food/g body weight) of ob/ob mice as a function of dose (log [g construct/kg BW]) using a dimer of the DhCpmFc(+)(Y349C)-GDF15(N3D):DhCpmFc(-)(S354C) heterodimer.

Figure 27 is a plot showing the effect on food intake (g food/g body weight) of ob/ob mice as a function of dose (log [g construct/kg BW]) using a dimer of the DhCpmFc(-)(Y349C)-GDF15:DhCpmFc(+)(S354C) heterodimer.

Figure 28 is a plot showing the effect on food intake (g food/g body weight) of ob/ob mice as a function of dose (log [g construct/kg BW]) using a dimer of the DhCpmFc(-)(Y349C)-GDF15(N3D):DhCpmFc(+)(S354C) heterodimer.

Figure 29 is a plot showing the effect on food intake (g food/g body weight) of ob/ob mice as a function of dose (log [g construct/kg BW]) using a dimer of the DhCpmFc(-)(Y349C)-GDF15(Ndel3):DhCpmFc(+)(S354C) heterodimer.

Figure 30 is a plot showing the effect on food intake (g food/g body weight) of ob/ob mice as a function of dose (log [g construct/kg BW]) using a dimer of the DhCpmFc(-)(Y349C)-G₄-GDF15(N3D):DhCpmFc(+)(S354C) heterodimer.

Figure 31 is a plot showing the effect on food intake (g food/g body weight) of ob/ob mice as a function of dose (log [g construct/kg BW]) using a dimer of the DhCpmFc(-)(Y349C)-(G₄S)₂-GDF15(N3D):DhCpmFc(+)(S354C) heterodimer.

Figure 32 is a plot showing the effect on food intake (g food/g body weight) of ob/ob mice as a function of dose (log [g construct/kg BW]) using a dimer of the DhCpmFc(-)(Y349C)-(G₄Q)₂-GDF15(N3D):DhCpmFc(+)(S354C) heterodimer.

Figure 33 is a plot showing the effect on food intake (g food/g body weight) of ob/ob mice as a function of dose (log [g construct/kg BW]) using a dimer of the DhCpmFc(-)(L351C)-(G₄S)₂-GDF15:DhCpmFc(+)(L351C) heterodimer.

Figure 34 is a plot showing the effect on food intake (g food/g body weight) of ob/ob mice as a function of dose (log [g construct/kg BW]) using a dimer of the HSA-GSAAQAAQQGS-GDF15 fusion protein.

Figure 35 is a plot showing the effect on food intake (g food/g body weight) of ob/ob mice as a function of dose (log [g construct/kg BW]) using a dimer of the HSA-GSPAPAPGS-GDF15 fusion protein.

Figure 36 is a plot showing the effect on food intake (g food/g body weight) of ob/ob mice as a function of dose (log [g construct/kg BW]) using a dimer of the HSA-GS(AAQAAQQ)₂GS-GDF15 fusion protein.

Figure 37 is a plot showing the effect on food intake (g food/g body weight) of ob/ob mice as a function of dose (log [g construct/kg BW]) using a dimer of the HSA-GS(PAPAP)₂GS-GDF15 fusion protein.

Figure 38 is a plot showing the effect on food intake (g food/g body weight) of ob/ob mice as a function of dose (log [g construct/kg BW]) using a dimer of the HSA-GDF15 fusion protein.

Figure 39 is a plot showing the effect on food intake (g food/g body weight) of ob/ob mice as a function of dose (log [g construct/kg BW]) using a dimer of the HSA-GGNAEAAAKEAAAKEAAKAGG-GDF15 fusion protein.

Figure 40 is a plot showing the effect on food intake (g food/g body weight) of ob/ob mice as a function of dose (log [g construct/kg BW]) using a dimer of the HSA-(G₄S)₆-GDF15 fusion protein.

Figure 41 is a plot showing the effect on food intake (g food/g body weight) of ob/ob mice as a function of dose (log [g construct/kg BW]) using a dimer of the DhCpmFc(-)(N297G)-GDF15(Ndel3):DhCpmFc(+)(N297G) heterodimer.

Figure 42 is a plot showing the effect on food intake (g food/g body weight) of ob/ob mice as a function of dose (log [g construct/kg BW]) using a dimer of the DhCpmFc(-)(N297G)-GDF15(N3D):DhCpmFc(+)(N297G) heterodimer.

Figure 43 is a plot showing the effect on food intake (g food/g body weight) of ob/ob mice as a function of dose (log [g construct/kg BW]) using a dimer of the DhCpmFc(-)(N297G)(Y349C)-GDF15(Ndel3):DhCpmFc(+)(N297G)(S354C) heterodimer.

Figure 44 is a plot showing the effect on food intake (g food/g body weight) of ob/ob mice as a function of dose (log [g construct/kg BW]) using a dimer of the DhCpmFc(-)(N297G)(Y349C)-GDF15(N3D):DhCpmFc(+)(N297G)(S354C) heterodimer.

Figure 45 is a plot showing the effect on food intake (g food/g body weight) of ob/ob mice as a function of dose (log [g construct/kg BW]) using a dimer of the DhCpmFc(-)(N297G)(L351C)-GDF15(Ndel3):DhCpmFc(+)(N297G)(L351C) heterodimer.

Figure 46 is a plot showing the effect on food intake (g food/g body weight) of ob/ob mice as a function of dose (log [g construct/kg BW]) using a dimer of the DhCpmFc(-)(N297G)(L351C)-GDF15(N3D):DhCpmFc(+)(N297G)(L351C) heterodimer.

Figure 47 is a plot showing the effect on food intake (g food/g body weight) of ob/ob mice as a function of dose (log [g construct/kg BW]) using a dimer of the DhCpmFc(-)(N297G)(A287C)-GDF15(Ndel3):DhCpmFc(+)(N297G)(L306C) heterodimer.

Figure 48 is a plot showing the effect on food intake (g food/g body weight) of ob/ob mice as a function of dose (log [g construct/kg BW]) using a dimer of the CpmFc(-)(N297G)-GDF15(Ndel3):CpmFc(+)(N297G) heterodimer.

Figure 49 is a plot showing the effect on food intake (g food/g body weight) of ob/ob mice as a function of dose (log [g construct/kg BW]) using a dimer of the Dh3CpmFc(-)-GDF15(Ndel3):Dh3CpmFc(N297G) heterodimer.

Figure 50 is a plot showing the effect on food intake (g food/g body weight) of ob/ob mice as a function of dose (log [g construct/kg BW]) using a dimer of the Dh3CpmFc(-)-GDF15(N3D):Dh3CpmFc(+)(N297G) heterodimer.

Figure 51 is a plot showing the effect on food intake (g food/g body weight) of ob/ob mice as a function of dose (log [g construct/kg BW]) using a dimer of the Dh3CpmFc(-)(Y349C)-GDF15(Ndel3):Dh3CpmFc(+)(N297G)(S354C) heterodimer.

Figure 52 is a plot showing the effect on food intake (g food/g body weight) of ob/ob mice as a function of dose (log [g construct/kg BW]) using a dimer of the Dh3CpmFc(-)(Y349C)-GDF15(N3D): Dh3CpmFc(-)(N297G)(S354C) heterodimer.

Figure 53 is a plot showing the effect on food intake (g food/g body weight) of ob/ob mice as a function of dose (log [g construct/kg BW]) using a dimer of the DhMonoFc(N297G)-GDF15 fusion protein.

Figure 54 is a plot of body weight (g) as a function of time (days post 1st injection) for vehicle, 0.1 nmol, 1 nmol and 10 nmol of a dimer of the Dh3CpmFc(-)-GDF15(N3D):Dh3CpmFc(+) heterodimer, and 0.1 nmol, 1 nmol and 10 nmol of a dimer of the Dh3ComFc(-)-GDF15(Ndel3) heterodimer.

Figure 55 is a bar graph of area under the curve (AUC) for a glucose tolerance test at week 2 of treatment with (a) vehicle (b) 10 nmol of a dimer of the Dh3CpmFc(-)-GDF15(N3D):Dh3CpmFc(+) heterodimer (c) 1 nmol of a dimer of the Dh3CpmFc(-)-

GDF15(N3D):Dh3CpmFc(+) heterodimer, (d) 0.1 mmol of a dimer of the Dh3CpmFc(-)-GDF15(N3D):Dh3CpmFc(+) heterodimer, (e) 10 mmol of a dimer of the Dh3CpmFc(-)-GDF15(Ndel3):Dh3CpmFc(+) heterodimer (g) 1 mmol of a dimer of the Dh3CpmFc(-)-GDF15(Ndel3):Dh3CpmFc(+) heterodimer, or (g) 0.1 mmol of a dimer of the Dh3CpmFc(-)-GDF15(Ndel):Dh3CpmFc(+) heterodimer.

Figure 56 is a bar graph of insulin (ng/mL) (fed) at week 3 of treatment with (a) vehicle (b) 10 mmol of a dimer of the Dh3CpmFc(-)-GDF15(N3D):Dh3CpmFc(+) heterodimer (c) 1 mmol of a dimer of the Dh3CpmFc(-)-GDF15(N3D):Dh3CpmFc(+) heterodimer, (d) 0.1 mmol of a dimer of the Dh3CpmFc(-)-GDF15(N3D):Dh3CpmFc(+) heterodimer, (e) 10 mmol of a dimer of the Dh3CpmFc(-)-GDF15(Ndel3):Dh3CpmFc(+) heterodimer (g) 1 mmol of a dimer of the Dh3CpmFc(-)-GDF15(Ndel3):Dh3CpmFc(+) heterodimer, or (g) 0.1 mmol of a dimer of the Dh3CpmFc(-)-GDF15(Ndel):Dh3CpmFc(+) heterodimer.

Figure 57 is a bar graph of triglycerides (mg/mL) (fed) at week 3 of treatment with (a) vehicle (b) 10 mmol of a dimer of the Dh3CpmFc(-)-GDF15(N3D):Dh3CpmFc(+) heterodimer (c) 1 mmol of a dimer of the Dh3CpmFc(-)-GDF15(N3D):Dh3CpmFc(+) heterodimer, (d) 0.1 mmol of a dimer of the Dh3CpmFc(-)-GDF15(N3D):Dh3CpmFc(+) heterodimer, (e) 10 mmol of a dimer of the Dh3CpmFc(-)-GDF15(Ndel3):Dh3CpmFc(+) heterodimer (g) 1 mmol of a dimer of the Dh3CpmFc(-)-GDF15(Ndel3):Dh3CpmFc(+) heterodimer, or (g) 0.1 mmol of a dimer of the Dh3CpmFc(-)-GDF15(Ndel):Dh3CpmFc(+) heterodimer.

Figure 58 is a bar graph of cholesterol (mg/mL) (fed) at week 3 of treatment with (a) vehicle (b) 10 mmol of a dimer of the Dh3CpmFc(-)-GDF15(N3D):Dh3CpmFc(+) heterodimer (c) 1 mmol of a dimer of the Dh3CpmFc(-)-GDF15(N3D):Dh3CpmFc(+) heterodimer, (d) 0.1 mmol of a dimer of the Dh3CpmFc(-)-GDF15(N3D):Dh3CpmFc(+) heterodimer, (e) 10 mmol of a dimer of the Dh3CpmFc(-)-GDF15(Ndel3):Dh3CpmFc(+) heterodimer (g) 1 mmol of a dimer of the Dh3CpmFc(-)-GDF15(Ndel3):Dh3CpmFc(+) heterodimer, or (g) 0.1 mmol of a dimer of the Dh3CpmFc(-)-GDF15(Ndel):Dh3CpmFc(+) heterodimer.

Figure 59 is a bar graph of area under the curve (AUC) for a glucose tolerance test at week 5 of treatment with (a) vehicle (b) 10 mmol of a dimer of the Dh3CpmFc(-)-GDF15(N3D):Dh3CpmFc(+) heterodimer (c) 1 mmol of a dimer of the Dh3CpmFc(-)-GDF15(N3D):Dh3CpmFc(+) heterodimer, (d) 0.1 mmol of a dimer of the Dh3CpmFc(-)-GDF15(N3D):Dh3CpmFc(+) heterodimer, (e) 10 mmol of a dimer of the Dh3CpmFc(-)-GDF15(Ndel3):Dh3CpmFc(+) heterodimer (g) 1 mmol of a dimer of the Dh3CpmFc(-)-GDF15(Ndel3):Dh3CpmFc(+) heterodimer, or (g) 0.1 mmol of a dimer of the Dh3CpmFc(-)-GDF15(Ndel):Dh3CpmFc(+) heterodimer.

Figure 60 is a bar graph showing SPR Binding (RU) with respect to FcγRI, FcγRIIA and FcγRIIA for (a) a dimer of DhCpmFc(-)-GDF15(Ndel3):DhCpmFc(+), (b) a dimer of DhCpmFc(-)(Y349C)-GDF15(Ndel3): DhCpmFc(+)(S354C); (c) a dimer of Dh3CpmFc(-)-GDF15(Ndel3); (d) a dimer of Dh3CpmFc(-)(Y349C)-GDF15(Ndel3)-Dh3CpmFc(+)(S354C); (e) a dimer of Dh3CpmFc(-)-GDF15(N3D); and (f) a dimer of Dh3CpmFc(-)(Y349C)-GDF15(N3D):Dh3CpmFc(+)(S354C) (from left to right with respect to each receptor).

Figure 61 is a bar graph showing DSC first Tm (°C) with respect to (a) (for pair of bars captioned “N3D”), a dimer of DhCpmFc(-)-GDF15(N3D):DhCpmFc(+) and a dimer of Dh3CpmFc(-)-GDF15(N3D):DhCpmFc(+); (b) (for the pair of bars captioned “Ndel3”), a dimer of DhCpmFc(-)-GDF15(Ndel3):DhCpmFc(+) and a dimer of Dh3CpmFc(-)-GDF15(Ndel3):DhCpmFc(+); (c) (for the pair of bars captioned “N3D+CC”), a dimer of DhCpmFc(-)(Y349C)-GDF15(N3D):DhCpmFc(+)(S354C) and a dimer of Dh3CpmFc(-)(Y349C)-GDF15(N3D):DhCpmFc(+)(S354C); and (d) (for the pair of bars captioned “Ndel3+CC”), a dimer of DhCpmFc(-)(Y349C)-GDF15(Ndel3):DhCpmFc(+)(S354C) and a dimer of Dh3CpmFc(-)(Y349C)-GDF15(Ndel3):DhCpmFc(+)(S354C).

DETAILED DESCRIPTION OF THE INVENTION

The instant disclosure provides fusion proteins comprising a GDF15 polypeptide or a GDF15 mutant polypeptides and constructs comprising such fusion proteins. Also provided is the generation and uses of the disclosed molecules, for example in treating a metabolic disorder, such as Type 2 diabetes, elevated glucose levels, elevated insulin levels, dyslipidemia or obesity. GDF15 polypeptides, GDF15 mutant polypeptides, and certain polypeptide constructs comprising GDF15 polypeptides and GDF15 mutant polypeptides, are described in co-owned PCT/US2012/032415, filed 04/05/2012, and PCT/2013/023465, filed 01/28/2013, both of which are expressly incorporated by reference herein for any purpose.

Recombinant polypeptide and nucleic acid methods used herein, including in the Examples, are generally those set forth in Sambrook et al., Molecular Cloning: A Laboratory Manual (Cold Spring Harbor Laboratory Press, 1989) or Current Protocols in Molecular Biology (Ausubel et al., eds., Green Publishers Inc. and Wiley and Sons 1994).

I. General Definitions

Following convention, as used herein “a” and “an” mean “one or more” unless specifically indicated otherwise.

As used herein, the terms “amino acid” and “residue” are interchangeable and, when used in the context of a peptide or polypeptide, refer to both naturally-occurring and synthetic

amino acids, as well as amino acid analogs, amino acid mimetics and non-naturally-occurring amino acids that are chemically similar to the naturally-occurring amino acids.

The terms “naturally-occurring amino acid” and “naturally encoded amino acid” are used interchangeably and refer to an amino acid that is encoded by the genetic code, as well as those amino acids that are encoded by the genetic code that are modified after synthesis, *e.g.*, hydroxyproline, γ -carboxyglutamate, and O-phosphoserine.

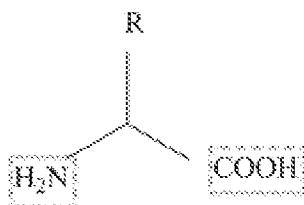
An “amino acid analog” is a compound that has the same basic chemical structure as a naturally-occurring amino acid, *i.e.*, an α carbon that is bound to a hydrogen, a carboxyl group, an amino group, and an R group, *e.g.*, homoserine, norleucine, methionine sulfoxide, methionine methyl sulfonium. Such analogs can have modified R groups (*e.g.*, norleucine) or modified peptide backbones, but will retain the same basic chemical structure as a naturally-occurring amino acid.

An “amino acid mimetic” is a chemical compound that has a structure that is different from the general chemical structure of an amino acid, but that functions in a manner similar to a naturally-occurring amino acid. Examples include a methacryloyl or acryloyl derivative of an amide, β -, γ -, δ -imino acids (such as piperidine-4-carboxylic acid) and the like.

The terms “non-naturally-occurring amino acid” and “non-naturally encoded amino acid” are used interchangeably and refer to a compound that has the same basic chemical structure as a naturally-occurring amino acid, but is not incorporated into a growing polypeptide chain by the translation complex. “Non-naturally-occurring amino acid” also includes, but is not limited to, amino acids that occur by modification (*e.g.*, posttranslational modifications) of a naturally encoded amino acid (including but not limited to, the 20 common amino acids) but are not themselves naturally incorporated into a growing polypeptide chain by the translation complex. A non-limiting lists of examples of non-naturally-occurring amino acids that can be inserted into a polypeptide sequence or substituted for a wild-type residue in polypeptide sequence include β -amino acids, homoamino acids, cyclic amino acids and amino acids with derivatized side chains. Examples include (in the L-form or D-form; abbreviated as in parentheses): citrulline (Cit), homocitrulline (hCit), $N\alpha$ -methylcitrulline (NMeCit), $N\alpha$ -methylhomocitrulline ($N\alpha$ -MeHoCit), ornithine (Orn), $N\alpha$ -Methylornithine ($N\alpha$ -MeOrn or NMeOrn), sarcosine (Sar), homolysine (hLys or hK), homoarginine (hArg or hR), homoglutamine (hQ), $N\alpha$ -methylarginine (NMeR), $N\alpha$ -methylleucine ($N\alpha$ -MeL or NMeL), N-methylhomolysine (NMeHoK), $N\alpha$ -methylglutamine (NMeQ), norleucine (Nle), norvaline (Nva), 1,2,3,4-tetrahydroisoquinoline (Tic), Octahydroindole-2-carboxylic acid (Oic), 3-(1-naphthyl)alanine (1-Nal), 3-(2-naphthyl)alanine (2-Nal), 1,2,3,4-tetrahydroisoquinoline (Tic), 2-indanylglycine (IgI), para-iodophenylalanine (pI-Phe), para-aminophenylalanine (4AmP or 4-Amino-Phe), 4-guanidino phenylalanine (Guf), glycyllsine (abbreviated “K(N ϵ -glycyl)” or “K(glycyl)” or

“K(gly)”), nitrophenylalanine (nitrophe), aminophenylalanine (aminophe or Amino-Phe), benzylphenylalanine (benzylphe), γ -carboxyglutamic acid (γ -carboxyglu), hydroxyproline (hydroxypro), p-carboxyl-phenylalanine (Cpa), α -aminoadipic acid (Aad), N α -methyl valine (NMeVal), N- α -methyl leucine (NMeLeu), N α -methylnorleucine (NMeNle),
 5 cyclopentylglycine (Cpg), cyclohexylglycine (Chg), acetylgarginine (acetylarg), α , β -diaminopropionioic acid (Dpr), α , γ -diaminobutyric acid (Dab), diaminopropionic acid (Dap), cyclohexylalanine (Cha), 4-methyl-phenylalanine (MePhe), β , β -diphenyl-alanine (BiPhA), aminobutyric acid (Abu), 4-phenyl-phenylalanine (or biphenylalanine; 4Bip), α -amino-isobutyric acid (Aib), beta-alanine, beta-aminopropionic acid, piperidinic acid, aminocaprioic
 10 acid, aminoheptanoic acid, aminopimelic acid, desmosine, diaminopimelic acid, N-ethylglycine, N-ethylasparagine, hydroxylysine, allo-hydroxylysine, isodesmosine, allo-isoleucine, N-methylglycine, N-methylisoleucine, N-methylvaline, 4-hydroxyproline (Hyp), γ -carboxyglutamate, ϵ -N,N,N-trimethyllysine, ϵ -N-acetyllysine, O-phosphoserine, N-acetylserine, N-formylmethionine, 3-methylhistidine, 5-hydroxylysine, ω -methylarginine, 4-
 15 Amino-O-Phthalic Acid (4APA), N-acetylglucosaminyl-L-serine, N-acetylglucosylaminyl-L-threonine, O-phosphotyrosine and other similar amino acids, and derivatized forms of any of those specifically listed.

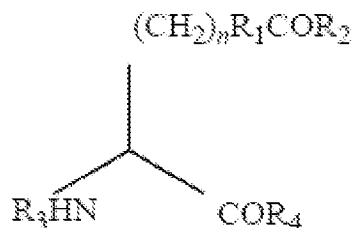
Also included in the definition of “non-naturally-occurring amino acid” is any amino acid comprising the structure



20

wherein the R group is any substituent other than the one used in the twenty natural amino acids.

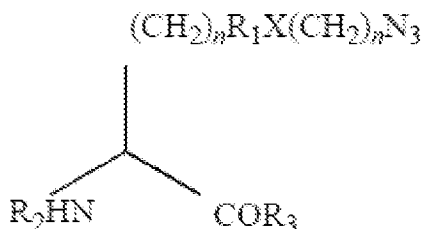
In some embodiments, the non-naturally encoded amino acid comprises a carbonyl group. In some embodiments, the non-naturally encoded amino acid has the structure:



wherein n is 0-10; R₁ is an alkyl, aryl, substituted alkyl, or substituted aryl; R₂ is H, an alkyl, aryl, substituted alkyl, and substituted aryl; and R₃ is H, an amino acid, a polypeptide, or an amino terminus modification group, and R₄ is H, an amino acid, a polypeptide, or a carboxy terminus modification group.

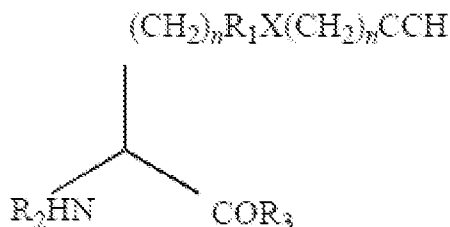
In some embodiments, the non-naturally encoded amino acid comprises an aminooxy group. In some embodiments, the non-naturally encoded amino acid comprises a hydrazide group. In some embodiments, the non-naturally encoded amino acid comprises a hydrazine group. In some embodiments, the non-naturally encoded amino acid residue comprises a semicarbazide group.

In some embodiments, the non-naturally encoded amino acid residue comprises an azide group. In some embodiments, the non-naturally encoded amino acid has the structure:



wherein n is 0-10; R₁ is an alkyl, aryl, substituted alkyl, substituted aryl or not present; X is O, N, S or not present; m is 0-10; R₂ is H, an amino acid, a polypeptide, or an amino terminus modification group, and R₃ is H, an amino acid, a polypeptide, or a carboxy terminus modification group.

In some embodiments, the non-naturally encoded amino acid comprises an alkyne group. In some embodiments, the non-naturally encoded amino acid has the structure:



wherein n is 0-10; R₁ is an alkyl, aryl, substituted alkyl, or substituted aryl; X is O, N, S or not present; m is 0-10, R₂ is H, an amino acid, a polypeptide, or an amino terminus modification group, and R₃ is H, an amino acid, a polypeptide, or a carboxy terminus modification group.

- 5 The term "substituted" means that a hydrogen atom on a molecule or group is replaced with a group or atom, which is referred to as a substituent. Typical substituents include: halogen, C₁₋₈alkyl, hydroxyl, C₁₋₈alkoxy, -NR^xR^x, nitro, cyano, halo or perhaloC₁₋₈alkyl, C₂₋₈alkenyl, C₂₋₈alkynyl, -SR^x, -S(=O)₂R^x, -C(=O)OR^x, -C(=O)R^x, wherein each R^x is independently hydrogen or C₁₋₈ alkyl. It is noted that when the substituent is -NR^xR^x, the
- 10 R^x groups may be joined together with the nitrogen atom to form a ring.

The term "alkyl" means a straight or branched chain hydrocarbon. Representative examples of alkyl groups include methyl, ethyl, propyl, isopropyl, butyl, isobutyl, tert-butyl, sec-butyl, pentyl and hexyl. Typical alkyl groups are alkyl groups having from 1 to 8 carbon atoms, which groups are commonly represented as C₁₋₈alkyl.

- 15 The term "alkoxy" means an alkyl group bonded to an oxygen atom. Representative examples of alkoxy groups include methoxy, ethoxy, tert-butoxy, propoxy and isobutoxy. Common alkoxy groups are C₁₋₈alkoxy.

The term "halogen" or "halo" means chlorine, fluorine, bromine or iodine.

- 20 The term "alkenyl" means a branched or straight chain hydrocarbon having one or more carbon-carbon double bonds. Representative examples alkenyl groups include ethenyl, propenyl, allyl, butenyl and 4-methylbutenyl. Common alkenyl groups are C₂₋₈alkenyl.

The term "alkynyl" means a branched or straight chain hydrocarbon having one or more carbon-carbon triple bonds. Representative examples of alkynyl groups include ethynyl, propynyl (propargyl) and butynyl. Common alkynyl groups are C₂₋₈alkynyl.

- 25 The term "cycloalkyl" means a cyclic, nonaromatic hydrocarbon. Examples of cycloalkyl groups include cyclopropyl, cyclobutyl, cyclopentyl, cyclohexyl and cycloheptyl. A cycloalkyl group can contain one or more double bond. Examples of cycloalkyl groups that contain double bonds include cyclopentenyl, cyclohexenyl, cyclohexadienyl and cyclobutadienyl. Common cycloalkyl groups are C₃₋₈ cycloalkyl groups.

The term "perfluoroalkyl" means an alkyl group in which all of the hydrogen atoms have been replaced with fluorine atoms. Common perfluoroalkyl groups are C₁₋₈perfluoroalkyl. An example of a common perfluoroalkyl group is -CF₃.

5 The term "acyl" means a group derived from an organic acid by removal of the hydroxy group (-OH). For example, the acyl group CH₃C(=O)- is formed by the removal of the hydroxy group from CH₃C(=O)OH.

The term "aryl" means a cyclic, aromatic hydrocarbon. Examples of aryl groups include phenyl and naphthyl. Common aryl groups are six to thirteen membered rings.

The term "heteroatom" as used herein means an oxygen, nitrogen or sulfur atom.

10 The term "heteroaryl" means a cyclic, aromatic hydrocarbon in which one or more carbon atoms of an aryl group have been replaced with a heteroatom. If the heteroaryl group contains more than one heteroatom, the heteroatoms may be the same or different. Examples of heteroaryl groups include pyridyl, pyrimidinyl, imidazolyl, thienyl, furyl, pyrazinyl, pyrrolyl, indolyl, triazolyl, pyridazinyl, indazolyl, purinyl, quinoliziny, isoquinolyl, quinolyl, 15 naphthyridinyl, quinoxalinyl, isothiazolyl and benzo[b]thienyl. Common heteroaryl groups are five to thirteen membered rings that contain from 1 to 4 heteroatoms. Heteroaryl groups that are five and six membered rings that contain 1 to 3 heteroatoms are particularly common.

The term "heterocycloalkyl" means a cycloalkyl group in which one or more of the carbon atoms has been replaced with a heteroatom. If the heterocycloalkyl group contains 20 more than one heteroatom, the heteroatoms may be the same or different. Examples of heterocycloalkyl groups include tetrahydrofuryl, morpholinyl, piperazinyl, piperidinyl and pyrrolidinyl. It is also possible for the heterocycloalkyl group to have one or more double bonds, but is not aromatic. Examples of heterocycloalkyl groups containing double bonds include dihydrofuran. Common heterocycloalkyl groups are three to ten membered rings 25 containing from 1 to 4 heteroatoms. Heterocycloalkyl groups that are five and six membered rings that contain 1 to 2 heteroatoms are particularly common.

It is also noted that the cyclic ring groups, i.e., aryl, heteroaryl, cycloalkyl, and heterocycloalkyl, can comprise more than one ring. For example, the naphthyl group is a fused bicyclic ring system. It is also intended that the present invention include ring groups 30 that have bridging atoms, or ring groups that have a spiro orientation.

Representative examples of five to six membered aromatic rings, optionally having one or two heteroatoms, are phenyl, furyl, thienyl, pyrrolyl, oxazolyl, thiazolyl, imidazolyl, pyrazolyl, isoxazolyl, isothiazolyl, pyridinyl, pyridiazinyl, pyrimidinyl, and pyrazinyl.

Representative examples of partially saturated, fully saturated or fully unsaturated 35 five to eight membered rings, optionally having one to three heteroatoms, are cyclopentyl, cyclohexyl, cycloheptyl, cyclooctyl and phenyl. Further exemplary five membered rings are furyl, thienyl, pyrrolyl, 2-pyrrolinyl, 3-pyrrolinyl, pyrrolidinyl, 1,3-dioxolanyl, oxazolyl,

thiazolyl, imidazolyl, 2H-imidazolyl, 2-imidazolynyl, imidazolidinyl, pyrazolyl, 2-pyrazolynyl, pyrazolidinyl, isoxazolyl, isothiazolyl, 1,2-dithiolyl, 1,3-dithiolyl, 3H-1,2-oxathiolyl, 1,2,3-oxadiazolyl, 1,2,4-oxadiazolyl, 1,2,5-oxadiazolyl, 1,3,4-oxadiazolyl, 1,2,3-triazolyl, 1,2,4-triazolyl, 1,3,4-thiadiazolyl, 3H-1,2,3-dioxazolyl, 1,2,4-dioxazolyl, 1,3,2-dioxazolyl, 1,3,4-dioxazolyl, 5H-1,2,5-oxathiazolyl, and 1,3-oxathiolyl.

Further exemplary six membered rings are 2H-pyranyl, 4H-pyranyl, pyridinyl, piperidinyl, 1,2-dioxinyl, 1,3-dioxinyl, 1,4-dioxanyl, morpholinyl, 1,4-dithianyl, thiomorpholinyl, pyndazinyl, pyrimidinyl, pyrazinyl, piperazinyl, 1,3,5-triazinyl, 1,2,4-triazinyl, 1,2,3-triazinyl, 1,3,5-trithianyl, 4H-1,2-oxazinyl, 2H-1,3-oxazinyl, 6H-1,3-oxazinyl, 6H-1,2-oxazinyl, 1,4-oxazinyl, 2H-1,2-oxazinyl, 4H-1,4-oxazinyl, 1,2,5-oxathiazinyl, 1,4-oxazinyl, o-isoxazinyl, p-isoxazinyl, 1,2,5-oxathiazinyl, 1,2,6-(3 oxathiazinyl, and 1,4,2-oxadiazinyl.

Further exemplary seven membered rings are azepinyl, oxepinyl, thiepinyl and 1,2,4-triazepinyl.

Further exemplary eight membered rings are cyclooctyl, cyclooctenyl and cyclooctadienyl.

Exemplary bicyclic rings consisting of two fused partially saturated, fully saturated or fully unsaturated five and/or six membered rings, optionally having one to four heteroatoms, are indolizinyl, indolyl, isoindolyl, indolinyl, cyclopenta(b)pyridinyl, pyrano(3,4-b)pyrrolyl, benzofuryl, isobenzofuryl, benzo(b)thienyl, benzo(c)thienyl, 1H-indazolyl, indoxazinyl, benzoxazolyl, anthranilyl, benzimidazolyl, benzthiazolyl, purinyl, quinolinyl, isoquinolinyl, cinnolinyl, phthalazinyl, quinazolinyl, quinoxalinyl, 1,8-naphthyridinyl, pteridinyl, indenyl, isoindenyl, naphthyl, tetralinyl, decalinyl, 2H-1-benzopyranyl, pyrido(3,4-b)pyridinyl, pyrido(3,2-b)pyridinyl, pyrido(4,3-b)-pyridinyl, 2H-1,3-benzoxazinyl, 2H-1,4-benzoxazinyl, 1H-2,3-benzoxazinyl, 4H-3,1-benzoxazinyl, 2H-1,2-benzoxazinyl and 4H-1,4-benzoxazinyl.

A cyclic ring group may be bonded to another group in more than one way. If no particular bonding arrangement is specified, then all possible arrangements are intended. For example, the term "pyridyl" includes 2-, 3-, or 4-pyridyl, and the term "thienyl" includes 2-, or 3-thienyl.

The term "isolated nucleic acid molecule" refers to a single or double-stranded polymer of deoxyribonucleotide or ribonucleotide bases read from the 5' to the 3' end (*e.g.*, a GDF15 nucleic acid sequence provided herein), or an analog thereof, that has been separated from at least about 50 percent of polypeptides, peptides, lipids, carbohydrates, polynucleotides or other materials with which the nucleic acid is naturally found when total nucleic acid is isolated from the source cells. Preferably, an isolated nucleic acid molecule is substantially free from any other contaminating nucleic acid molecules or other molecules

that are found in the natural environment of the nucleic acid that would interfere with its use in polypeptide production or its therapeutic, diagnostic, prophylactic or research use.

The term “isolated polypeptide” refers to a polypeptide (*e.g.*, a GDF15 polypeptide or GDF15 mutant polypeptide provided herein) that has been separated from at least about 50 percent of polypeptides, peptides, lipids, carbohydrates, polynucleotides, or other materials with which the polypeptide is naturally found when isolated from a source cell. Preferably, the isolated polypeptide is substantially free from any other contaminating polypeptides or other contaminants that are found in its natural environment that would interfere with its therapeutic, diagnostic, prophylactic or research use.

The term “encoding” refers to a polynucleotide sequence encoding one or more amino acids. The term does not require a start or stop codon. An amino acid sequence can be encoded in any one of the different reading frames provided by a polynucleotide sequence.

The terms “identical” and percent “identity,” in the context of two or more nucleic acids or polypeptide sequences, refer to two or more sequences or subsequences that are the same. “Percent identity” means the percent of identical residues between the amino acids or nucleotides in the compared molecules and is calculated based on the size of the smallest of the molecules being compared. For these calculations, gaps in alignments (if any) can be addressed by a particular mathematical model or computer program (*i.e.*, an “algorithm”). Methods that can be used to calculate the identity of the aligned nucleic acids or polypeptides include those described in *Computational Molecular Biology*, (Lesk, A. M., ed.), (1988) New York: Oxford University Press; *Biocomputing Informatics and Genome Projects*, (Smith, D. W., ed.), 1993, New York: Academic Press; *Computer Analysis of Sequence Data, Part I*, (Griffin, A. M., and Griffin, H. G., eds.), 1994, New Jersey: Humana Press; von Heinje, G., (1987) *Sequence Analysis in Molecular Biology*, New York: Academic Press; *Sequence Analysis Primer*, (Gribskov, M. and Devereux, J., eds.), 1991, New York: M. Stockton Press; and Carillo *et al.*, (1988) *SIAM J. Applied Math.* 48:1073.

In calculating percent identity, the sequences being compared are aligned in a way that gives the largest match between the sequences. The computer program used to determine percent identity is the GCG program package, which includes GAP (Devereux *et al.*, (1984) *Nucl. Acid Res.* 12:387; Genetics Computer Group, University of Wisconsin, Madison, WI). The computer algorithm GAP is used to align the two polypeptides or polynucleotides for which the percent sequence identity is to be determined. The sequences are aligned for optimal matching of their respective amino acid or nucleotide (the “matched span”, as determined by the algorithm). A gap opening penalty (which is calculated as 3x the average diagonal, wherein the “average diagonal” is the average of the diagonal of the comparison matrix being used; the “diagonal” is the score or number assigned to each perfect amino acid match by the particular comparison matrix) and a gap extension penalty (which is usually

1/10 times the gap opening penalty), as well as a comparison matrix such as PAM 250 or BLOSUM 62 are used in conjunction with the algorithm. In certain embodiments, a standard comparison matrix (*see*, Dayhoff *et al.*, (1978) *Atlas of Protein Sequence and Structure* 5:345-352 for the PAM 250 comparison matrix; Henikoff *et al.*, (1992) *Proc. Natl. Acad. Sci. U.S.A.* 89:10915-10919 for the BLOSUM 62 comparison matrix) is also used by the algorithm.

Recommended parameters for determining percent identity for polypeptides or nucleotide sequences using the GAP program are the following:

- Algorithm: Needleman *et al.*, 1970, *J. Mol. Biol.* 48:443-13;
- 10 Comparison matrix: BLOSUM 62 from Henikoff *et al.*, 1992, *supra*;
- Gap Penalty: 12 (but with no penalty for end gaps)
- Gap Length Penalty: 4
- Threshold of Similarity: 0

Certain alignment schemes for aligning two amino acid sequences can result in matching of only a short region of the two sequences, and this small aligned region can have very high sequence identity even though there is no significant relationship between the two full-length sequences. Accordingly, the selected alignment method (*e.g.*, the GAP program) can be adjusted if so desired to result in an alignment that spans at least 50 contiguous amino acids of the target polypeptide.

The term "GDF15 polypeptide" means a naturally-occurring or "wild-type" GDF15 expressed in a mammal, including without limitation, a human, rabbit, monkey (*e.g.* cynomolgous monkey), dog, rat mouse or pig. In one aspect, GDF15 polypeptide refers to any full-length GDF15, *e.g.*, SEQ ID NO:4, which consists of 308 amino acid residues and which is encoded by the nucleotide sequence SEQ ID NO:3; any form comprising the active and prod domains of the polypeptide, *e.g.*, SEQ ID NO:8, which consists of 279 amino acid residues and which is encoded by the nucleotide sequence SEQ ID NO:7, and in which the 29 amino acid residues at the amino-terminal end of the full-length GDF15 (*i.e.*, which constitute the signal peptide) have been removed; and any form of GDF15 comprising the active domain from which the prodomain and signal sequence have been removed, *e.g.*, SEQ ID NO:12, which consists of 112 amino acid residues and which is encoded by the nucleotide sequence SEQ ID NO:11, in which the signal sequence and the prodomain have been removed. GDF15 polypeptides can but need not comprise an amino-terminal methionine, which may be introduced by engineering or as a result of a bacterial expression process.

The term "GDF15 mutant polypeptide" refers to a GDF15 polypeptide (*e.g.*, SEQ ID NOs:4, 8 or 12) that has been modified. Such modifications include, but are not limited to, one or more amino acid substitutions, including substitutions with non-naturally-occurring amino acids non-naturally-occurring amino acid analogs and amino acid mimetics, additions

or deletions. In one aspect, the term “GDF15 mutant polypeptide” refers to a GDF15 polypeptide in which at least one residue normally found at a given position of the naturally-occurring GDF15 polypeptide is deleted or is replaced by a residue not normally found at that position in the naturally-occurring GDF15 polypeptide. In some cases it will be desirable to replace a single residue normally found at a given position in the sequence of a naturally-occurring GDF15 polypeptide with more than one residue that is not normally found at the position; in still other cases it may be desirable to maintain the sequence of the naturally-occurring GDF15 polypeptide and insert one or more residues at a given position in the protein; in still other cases it may be desirable to delete a given residue entirely; all of these constructs are encompassed by the term “GDF15 mutant polypeptide.”

In various embodiments, a GDF15 mutant polypeptide comprises an amino acid sequence that is at least about 85 percent identical to a naturally-occurring GDF15 polypeptide (*e.g.*, SEQ ID NOs:4, 8 or 12). In other embodiments, a GDF15 mutant polypeptide comprises an amino acid sequence that is at least about 90 percent, or about 95, 96, 97, 98, or 99 percent identical to a naturally-occurring GDF15 polypeptide amino acid sequence (*e.g.*, SEQ ID NOs:4, 8 or 12). Such GDF15 mutant polypeptides preferably, but need not, possess at least one activity of a naturally-occurring GDF15 polypeptide, such as the ability to lower blood glucose, insulin, triglyceride, or cholesterol levels; the ability to reduce body weight; the ability to improve glucose tolerance, lipid tolerance, or insulin sensitivity; or the ability to lower urine glucose and protein excretion.

The term “GDF15 region” encompasses GDF15 polypeptides and GDF15 mutant polypeptides and sequences as defined above.

A GDF15 polypeptide or a GDF15 mutant polypeptide is preferably biologically active. In some instances, a GDF15 polypeptide or a GDF15 mutant polypeptide used to treat or ameliorate a metabolic disorder in a subject from or derived from a different species as the subject. In some instances, a GDF15 polypeptide or a GDF15 mutant polypeptide used to treat or ameliorate a metabolic disorder in a subject from or derived from the same species as the subject.

The term “native Fc” refers to molecule or sequence comprising the sequence of a non-antigen-binding fragment resulting from digestion of whole antibody, whether in monomeric or multimeric form. The original immunoglobulin source of the native Fc is preferably of human origin and may be any of the immunoglobulins, although IgG1 and IgG2 are preferred. Native Fc's are made up of monomeric polypeptides that may be linked into dimeric or multimeric forms by covalent (*i.e.*, disulfide bonds) and non-covalent association. The number of intermolecular disulfide bonds between monomeric subunits of native Fc molecules ranges from 1 to 4 depending on class (*e.g.*, IgG, IgA, IgE) or subclass (*e.g.*, IgG1, IgG2, IgG3, IgA1, IgGA2). One example of a native Fc is a disulfide-bonded dimer resulting

from papain digestion of an IgG (see Ellison et al. (1982), *Nucleic Acids Res.* 10: 4071-9). The term "native Fc" as used herein is generic to the monomeric, dimeric, and multimeric forms.

The term "Fc variant" refers to a molecule or sequence that is modified from a native Fc. Such modifications include, but are not limited to, one or more amino acid substitutions, including substitutions with non-naturally-occurring amino acids, non-naturally-occurring amino acid analogs and amino acid mimetics, deletions or additions. Thus, the term "Fc variant" comprises a molecule or sequence that is humanized from a non-human native Fc. Furthermore, a native Fc comprises sites that may be removed because they provide structural features or biological activity that are not required for the fusion molecules of the present invention. Thus, the term "Fc variant" comprises a molecule or sequence that lacks one or more native Fc sites or residues that affect or are involved in (1) disulfide bond formation, (2) incompatibility with a selected host cell (3) N-terminal heterogeneity upon expression in a selected host cell, (4) glycosylation, (5) interaction with complement, (6) binding to an Fc receptor other than a salvage receptor, or (7) antibody-dependent cellular cytotoxicity (ADCC). Fc variants are described in further detail hereinafter. In various embodiments, an Fc variant comprises an amino acid sequence that is at least about 85 percent identical to a native Fc. In other embodiments, an Fc variant comprises an amino acid sequence that is at least about 90 percent, or about 95, 96, 97, 98, or 99 percent identical to a native Fc.

The term "Fc domain" encompasses native Fc and Fc variant molecules and sequences as defined above. As with Fc variants and native Fcs, the term "Fc domain" includes molecules in monomeric or multimeric form.

The term "multimer" as applied to Fc domains or molecules comprising Fc domains refers to molecules having two or more polypeptide chains associated covalently, noncovalently, or by both covalent and non-covalent interactions. IgG molecules typically form dimers; IgM, pentamers; IgD, dimers; and IgA, monomers, dimers, trimers, or tetramers. Multimers may be formed by exploiting the sequence and resulting activity of the native Ig source of the Fc or by derivatizing such a native Fc.

The term "dimer" as applied to Fc domains or molecules comprising Fc domains refers to molecules having two polypeptide chains associated covalently, non-covalently or by both covalent and non-covalent interactions.

The term "hinge" or "hinge region" herein includes the flexible polypeptide comprising the amino acids between the first and second constant domains of an antibody. The "hinge region" as referred to herein is a sequence region of 6-62 amino acids in length, only present in IgA, IgD and IgG, which encompasses the cysteine residues that bridge the two heavy chains.

The “polypeptide linker” refers to a short polypeptide, generally from 1 to 30 amino acid residues in length, that covalently links together two polypeptides, typically via peptide bonds.

As used herein, the term HSA polypeptide encompasses a naturally-occurring “wild type” human serum albumin. The term also includes various bioactive fragments and variants, fusion proteins, and modified forms of the wild type HSA protein. Such bioactive fragments or variants, fusion proteins, and modified forms of wild type HSA protein have at least a portion of the amino acid sequence of substantial sequence identity to the wild type HAS protein. In certain embodiments, such bioactive fragments or variants, fusion proteins, and modified forms of wild type HSA protein have an amino acid sequence that is at least about 85 percent identical to the wild type HSA. In other embodiments, such bioactive fragments or variants, fusion proteins, and modified forms of wild type HSA protein have an amino acid sequence that is at least about 90 percent, or about 95, 96, 97, 98, or 99 percent identical to the wild type HAS.

II. Fc Fusions Comprising GDF15 Polypeptides or GDF15 Mutant Polypeptides, and Polynucleotides

A range of Fc fusion proteins comprising a GDF15 polypeptide or a GDF15 mutant polypeptide are provided herein. In some embodiments, the fusion proteins comprise a native Fc. In some embodiments, the fusion proteins comprise an Fc domain that has been engineered.

In some embodiments, the GDF15 polypeptide- (or GDF15 mutant polypeptide-) containing Fc fusion proteins associate with another polypeptide chain consisting of or comprising an Fc domain to form a heterodimer. In some embodiments, two such heterodimers associate to form a heterotetramer. Some of these polypeptide constructs (i.e., GDF15 polypeptide- (or GDF15 mutant polypeptide-) containing Fc fusion proteins and multimers comprising one or more such GDF15 polypeptide- (or GDF15 mutant polypeptide-) containing Fc fusion proteins) were studied empirically, as described in the Examples presented herein below.

Antibodies belong to the immunoglobulin class of proteins which includes IgG, IgA, IgE, IgM, and IgD. The most abundant immunoglobulin class in human serum is IgG (Deisenhofer J 1981 *Biochem* 20:2361-2370; Huber R 1984 *Behring Inst Mitt* 76:1-14; Roux KH 1999 *Int Arch Allergy Immunol* 120:85-99). The IgG structure has four chains, two light and two heavy chains; each light chain has two domains and each heavy chain has four domains. The antigen binding site is located in the Fab domain (Fragment antigen binding) which contains a variable light (VL) and a variable heavy (VH) chain domain as well as constant light (LC) and constant heavy (CH1) chain domains. The CH2 and CH3 domain

region of the heavy chain is called Fc (Fragment crystallizable). The IgG molecule can be considered as a heterotetramer having two heavy chains that are held together by disulfide bonds (-S-S-) at the hinge region and two light chains. The number of hinge disulfide bonds varies among the immunoglobulin subclasses (Papadea C 1989 *Crit Rev Clin Lab Sci* 27:27-58). The FcRn binding site is located in the Fc domain of the antibody (Martin WL 2001 *Mol Cell* 7:867-877), and thus the extended serum half-life property of the antibody is retained in the Fc fragment. The Fc domain alone can be thought of as a homodimer of heavy chains comprising CH2 and CH3 domains.

In certain preferred embodiments, the fusion proteins described herein comprise an IgG Fc domain, derived from a wild-type human IgG Fc domain. By "wild-type" human IgG Fc, it is meant a sequence of amino acids that occurs naturally within the human population. Of course, just as the Fc sequences may vary slightly between individuals, one or more alterations may be made to a wild-type sequence and still remain within the scope of the invention. For example, the Fc domain may contain additional alterations that are not related to the present invention, such as a mutation in a glycosylation site or the inclusion of an unnatural amino acid. In certain embodiments, the polypeptide containing the CH3 region is an IgG molecule and further contains a CH1 and CH2 domain. Exemplary human IgG sequences comprise the constant regions of IgG1, IgG2, IgG3 and IgG4. The Fc domain also may be comprised within the constant region of an IgA, IgD, IgE, and IgM heavy chain.

Some Fc fusion proteins containing GDF15 polypeptide or GDF15 mutant polypeptide, and multimers comprising such Fc fusion proteins include those described below.

II.A. DhMonoFc Constructs

The designations "Mono-" or "MonoFc domain" in the instant disclosure refer to an Fc domain that has been engineered to reduce or prevent the formation of homodimers. In one embodiment, a MonoFc domain is provided by introducing a tyrosine to threonine mutation (Y349T) and two lysine to aspartic acid mutations (K392D and K409D) in a native Fc or Fc variant.

The C-terminal lysine (K447) optionally may be deleted in the MonoFc. This may be advantageous, for example, when a peptide is fused at the C-terminus to reduce proteolysis of the fusion protein.

A fusion protein is provided comprising a MonoFc domain and a GDF15 region. Typically, the N-terminus of the GDF15 region is linked, directly or via a polypeptide linker, to the C-terminus of the MonoFc domain. However, in some embodiments, the N-terminus of the MonoFc domain is linked, directly or via a polypeptide linker, to the C-terminus of the GDF15 region.

In certain embodiments, a dimer is provided comprising two such fusion proteins linked via an interchain disulfide bond between their respective GDF15 regions. In some embodiments the dimer is a homodimer. In other embodiments, the dimer is a heterodimer.

The designations “DhMono-” or “DhMonoFc domain” in the instant disclosure refer to an Fc domain from which all or part of the hinge region has been removed that has been engineered to reduce or prevent the formation of homodimers. In one embodiment, a DhMonoFc domain is provided introducing a tyrosine to threonine mutation (Y349T) and two lysine to aspartic acid mutations (K392D and K409D) in a native Fc or Fc variant from which all or part of the hinge region has been removed.

The C-terminal lysine (K447) optionally may be deleted in the DhMonoFc. This may be advantageous, for example, when a peptide is fused at the C-terminus to reduce proteolysis of the fusion protein.

A fusion protein is provided comprising a DhMonoFc domain and a GDF15 region. Typically, the N-terminus of the GDF15 region is linked, directly or via a polypeptide linker, to the C-terminus of the DhMonoFc domain. However, in some embodiments, the N-terminus of the DhMonoFc domain is linked, directly or via a polypeptide linker, to the C-terminus of the GDF15 region.

In certain embodiments, a dimer is provided comprising two such fusion proteins linked via an interchain disulfide bond between their respective GDF15 regions. In some embodiments the dimer is a homodimer. In other embodiments, the dimer is a heterodimer.

II.A.1 DhMonoFc-GDF15

The designation “MonoFc-GDF15” in the instant disclosure refers to a fusion protein comprising a GDF15 polypeptide, the N-terminus of which is linked directly to the C-terminus of a MonoFc domain.

In certain embodiments, a homodimer is provided comprising two such fusion proteins linked via an interchain disulfide bond between their respective GDF15 regions.

More particularly, in a specific embodiment, the homodimer comprises:

(a) two MonoFc domains (one each monomer) comprising the sequence:

30 APELLGGPSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVD
GVEVHNAKTKPREEQYNSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPA
PIEKTISKAKGQPREPQVTTLPSSREEMTKNQVSLTCLVKGFYPSDIAVE
WESNGQPENNYDTTPVLDSDGSFFLYSDLTVDKSRWQQGNVVFSCSVME
ALHNHYTQKSLSLSPG (SEQ ID NO:22),

35 and

(b) two GDF15 polypeptides (one each monomer) comprising the sequence of SEQ ID NO:12.

In a preferred embodiment, the fusion protein comprises the amino acid sequence:

APELLGGPSVFLFPPKPKDTLMI SRTPEVTCVVVDVSHEDPEVKFNWYVD
 5 GVEVHNAKTKPREEQYNSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPA
 PIEKTISKAKGQPREPQVTTLPSSREEMTKNQVSLTCLVKGFYPSDIAVE
 WESNGQPENNYDTPPVLDSDGSFFLYSDLTVDKSRWQQGNVFSQSVSMHE
 ALHNHYTQKSLSLSPGARNGDHCPLGPGRCCRLHTVRASLEDLGWADWVL
 SPREVQVTMCIGACPSQFRAANMHAQIKTSLHRLKPDTPAPCCVPASYN
 10 PMVLIQKTDGTGVSLSQTYDDLLAKDCHCI (SEQ ID NO:46),

which is encoded by the nucleic acid sequence:

gcacctgaactcctggggggaccgtcagtccttcctcttccccc
 caaaacccaaggacaccctcatgatctcccgacccctgaggtcacatgc
 gtggtggtggacgtgagccacgaagaccctgaggtcaagttcaactggta
 15 cgtggacggcgtggaggtgcataatgccaagacaaagccgcgggaggagc
 agtacaacagcacgtaccgtgtggtcagcgtcctcacctcctgcaccag
 gactggctgaatggcaaggagtacaagtgaaggtctccaacaaagccct
 cccagcccccatcgagaaaaccatctccaaagccaaagggcagccccgag
 aaccacaggtgaccaccctgcccccatcccgaggagatgaccaagaac
 20 caggtcagcctgacctgctggtcaaaggcttctatcccagcgacatcgc
 cgtggagtgaggagagcaatgggcagccggagaaactacgacaccacgc
 ctcccgctgctggactccgacggctccttcttctcctctatagcgacctcacc
 gtggacaagagcaggtggcagcaggggaacgtcttctcatgctccgtgat
 gcatgaggctctgcacaaccactacacgcagaagagcctctccctgtctc
 25 cgggtgcgcgcaacggagaccactgtccgctcgggccccggcggttgctgc
 cgtctgcacacgggtccgcgcgtcgttggaagacctgggctgggcccattg
 ggtgctgtcgccacgggaggtgcaagtgacctgtgcatcggcgcgtgcc
 cgagccagttccgggaggaaacatgcacgcgcagatcaagacgagcctg
 caccgcctgaagcccgcacacgggtgccagcgcctgctgcgtgcccgcag
 30 ctacaatcccattggtgctcattcaaaagaccgacaccggggtgtcgtcc
 agacctatgatgacttgtagccaaagactgccactgcata (SEQ ID NO:45).

As discussed above, in a specific embodiment, a homodimer is provided comprising two monomers having the sequence of SEQ ID NO:46.

35 II.A.2 *DhMonoFc-(G₄S)₄-GDF15*

The designation "DhMonoFc-(G₄S)₄-GDF15" in the instant disclosure refers to a fusion protein comprising a GDF15 polypeptide linked to a DhMonoFc domain via a polypeptide linker comprising the sequence of SEQ ID NO:18 that connects the N-terminus of the GDF15 polypeptide to the C-terminus of the DhMonoFc domain.

5 In certain embodiments, a homodimer is provided comprising two such fusion proteins linked via an interchain disulfide bond between their respective GDF15 regions.

More particularly, in a specific embodiment, the homodimer comprises:

(a) two DhMonoFc domains (one each monomer) comprising the sequence of SEQ ID NO:22,

10 (b) two GDF15 polypeptides (one each monomer) comprising the sequence of SEQ ID NO:12, and

(c) two polypeptide linkers (one each monomer) comprising the sequence:

GGGGSGGGSGGGSGGGGS (SEQ ID NO:18)

each linking the N-terminus of a GDF15 polypeptide to the C-terminus of a DhMonoFc domain.

15 In a preferred embodiment, the fusion protein comprises the amino acid sequence (linker sequence double underlined):

APELLGGPSVFLFPPKPKDTLMISRTPEVTCVVDVSHEDPEVKFNWYVD
 GVEVHNAKTKPREEQYNSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPA
 20 PIEKTIKAKGQPREPQVTTLPSSREEMTKNQVSLTCLVKGFYPSDIAVE
 WESNGQPENNYDTPPVLDSDGSFFLYSDLTVDKSRWQQGNVFSQSVME
 ALHNHYTQKSLSLSPGGGGSGGGSGGGSGGGGSARNGDHCPLGPGRC
 CRLHTVRASLEDLGWADWVLSPREVQVTCIGACPSQFRAANMHAQIKTS
 LHRLKPDTVPAPCCVPASYNPMVLIQKTDGTGVSLLQTYDDLLAKDCHCI (SEQ ID
 25 NO:24),

which is encoded by the nucleic acid sequence:

cacctgaactcctggggggacgctcagtccttcttccccccaaaaccc
 aaggacaccctcatgatctcccggaaccctgaggtcacatgcgtggtggt
 ggacgtgagccacgaagaccctgaggtcaagttcaactggtacgtggacg
 30 gcgtggaggtgcataatgccaagacaaagccgcggaggagcagtacaac
 agcacgtaccgtgtggtcagcgtcctcaccgtcctgcaccaggactggct
 gaatggcaaggagtacaagtgaaggtctccaacaaagccctcccagccc
 ccatcgagaaaaccatctccaaagccaaaggcagccccgagaaccacag
 gtgaccaccctgccccatcccgaggagatgaccaagaaccagggtcag

cctgacctgcctgggtcaaaggcttctatcccagcgacatcgccgtggagt
 gggagagcaatgggcagccggagaacaactacgacaccacgcctcccgtg
 ctggactccgacggctccttcttctctatagcgacctcaccgtggacaa
 gagcaggtggcagcaggggaacgtcttctcatgctccgtgatgcatgagg
 5 ctctgcacaaccactacacgcagaagagcctctccctgtctccgggtgga
 ggtggtggatccggaggcgggtggaagcggaggtggtggatctggaggcgg
 tgggaagcgcgcgaacggagaccactgtccgctcgggccccgggcgttgct
 gccgtctgcacacggtccgcgcgtcgtggaagacctgggctgggcccgat
 tgggtgctgtcgccacgggaggtgcaagtgaccatgtgcatcggcgcgtg
 10 cccgagccagttccgggaggcaaacatgcacgcgcagatcaagacgagcc
 tgcaccgcctgaagcccgacacgggtgccagcgccctgctgctgcccgcc
 agctacaatcccagtggtgctcattcaaaagaccgacaccggggtgctcgt
 ccagacctatgatgacttgtagccaaagactgccactgcatatga (SEQ ID NO:23) .

As discussed above, in a specific embodiment, a homodimer is provided comprising
 15 two monomers having the sequence of SEQ ID NO:24.

II.A.3 DhMonoFc-(G₄S)₄-GDF15(H6D)

The designation “DhMonoFc-(G₄S)₄-GDF15(H6D)” in the instant disclosure refers to
 fusion protein comprising a GDF15 polypeptide linked to a DhMonoFc domain via a
 20 polypeptide linker comprising the sequence of SEQ ID NO:18 that connects the N-terminus
 of the GDF15 polypeptide to the C-terminus of the DhMonoFc domain. The GDF15(H6D)
 variant is a naturally-occurring human GDF15 variant.

In certain embodiments, a homodimer is provided comprising two such fusion
 proteins linked via an interchain disulfide bond between their respective GDF15 regions.

25 More particularly, in a specific embodiment, the homodimer comprises:

(a) two DhMonoFc domains (one each monomer) comprising the sequence of SEQ
 ID NO:22,

(b) two GDF15(H6D) polypeptides (one each monomer) comprising the sequence:

ARNGDDCPLGPGRCRLHTVRASLEDLGWADWVLSPREVQVTMCIGACPS
 30 QFRAANMHAQIKTSLHRLKPDTPAPCCVPASYNPMVLIQKTDGTGVSLOT
 YDDLAKDCHCI (SEQ ID NO:25) ,

and

(c) two polypeptide linkers (one each monomer) comprising the sequence of SEQ ID
 NO:18 each linking the N-terminus of the GDF15 polypeptide to the C-terminus of the
 35 DhMonoFc domain.

In a preferred embodiment, the fusion protein comprises the amino acid sequence (linker sequence double underlined):

APELLGGPSVFLFPPKPKDTLMISRTPEVTCVVDVSHEDPEVKFNWYVD
 GVEVHNAKTKPREEQYNSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPA
 5 PIEKTISKAKGQPREPQVTTLPSSREEMTKNQVSLTCLVKGFYPSDIAVE
 WESNGQPENNYDTTPPVLDSDGSFFLYSDLTVDKSRWQQGNVFSQSVME
 ALHNHYTQKSLSLSPGGGGSGGGSGGGSGGGSGGGGSARNGDDCPLGPGRC
 CRLHTVRASLEDLGWADWVLSPREVQVTMCIGACPSQFRAANMHAQIKTS
 LHRLKPDTPVAPCCVPASYNPMVLIQKTDTGVSLSQTYDDLLAKDCHCI (SEQ ID
 10 NO:27),

which is encoded by the nucleic acid sequence:

gcacctgaactcctggggggaccgtcagtccttctctccccccaaaacc
 caaggacaccctcatgatctccggaccctgaggtcacatgcgtggtgg
 tggacgtgagccacgaagaccctgaggtcaagttcaactggtacgtggac
 15 ggcgtggaggtgcataatgccaaagacaaagccgcgggaggagcagtacaa
 cagcacgtaccgtgtggtcagcgtcctcaccgtcctgcaccaggactggc
 tgaatggcaaggagtacaagtgaaggtctccaacaaagccctcccagcc
 cccatcgagaaaaccatctccaaagccaaagggcagccccgagaaccaca
 ggtgaccaccctgcccccatcccgaggagatgaccaagaaccaggtca
 20 gcctgacctgcctggtcaaaggcttctatcccagcgacatcgccgtggag
 tgggagagcaatgggcagccggagaacaactacgacaccacgcctcccgt
 gctggactccgacggtccttctctctctatagcgacctcaccgtggaca
 agagcaggtggcagcaggggaacgtcttctcatgctccgtgatgcatgag
 gctctgcacaaccactacacgcagaagagcctctccctgtctccgggtgg
 25 aggtggtggatccggaggcgggtggaagcggaggtggtggatctggaggcg
 gtggaagcgcgcgcaacggagacgactgtccgctcgggcccgggcgttgc
 tgccgtctgcacacgggtccgcgcgtcgctggaagacctgggctgggcca
 ttgggtgctgtcgccacgggaggtgcaagtgacctgtgcatcggcgcgt
 gcccagaccagttccgggcccgaacatgcacgcgcagatcaagacgagc
 30 ctgcaccgcctgaagcccgacacgggtgccagcgccctgctgcgtgcccgc
 cagctacaatcccattggtgctcattcaaaagaccgacaccggggtgtgcg
 tccagacctatgatgacttgttagccaaagactgccactgcatatga (SEQ ID
 NO:26).

As discussed above, in a specific embodiment, a homodimer is provided comprising
 35 two monomers having the sequence of SEQ ID NO:27.

II.B. HemiFc

The designations “Hemi-” or “HemiFc domain” in the instant disclosure refer to a polypeptide chain comprising a first Fc domain linked, directly or via a polypeptide linker, to a second Fc domain. In one embodiment, the first Fc domain and the second Fc domain have the same sequence. In another embodiment, the first Fc domain and second Fc domain have different sequences. Typically, the first and second Fc domains are covalently associated via disulfide bonds between their respective hinge regions.

The C-terminal lysine (K447) optionally may be deleted in the first Fc domain, the second Fc domain, or both. This may be advantageous, for example, when a peptide is fused at the C-terminus to reduce proteolysis of the fusion protein.

A fusion protein is provided comprising a HemiFc domain and a GDF15 region. Typically, the N-terminus of the GDF15 region is linked, directly or via a polypeptide linker, to the C-terminus of the HemiFc domain. However, in some embodiments, the N-terminus of the HemiFc domain is linked, directly or via a polypeptide linker, to the C-terminus of the GDF15 region.

In certain embodiments, a dimer is provided comprising two such fusion proteins linked via an interchain disulfide bond between their respective GDF15 regions. In some embodiments the dimer is a homodimer. In other embodiments, the dimer is a heterodimer. See Figure 3 for a graphic depiction of an embodiment of a homodimer of two GGG-Fc-(G₄S)₄-Fc-S(G₄S)₄-GDF15 fusion proteins.

GGGFc-(G₄S)₄-Fc-S(G₄S)₄-GDF15

The designation “GGGFc-(G₄S)₄-Fc-S(G₄S)₄-GDF15” in the instant disclosure refers to a fusion protein comprising a GDF15 polypeptide linked to HemiFc domain via a first polypeptide linker comprising the sequence of SEQ ID NO:30, that connects the N-terminus of the GDF15 polypeptide to the C-terminus of the HemiFc domain. The HemiFc domain comprises a first Fc domain linked to a second Fc domain via a polypeptide linker comprising the sequence of SEQ ID NO:18, that connects the N-terminus of the first Fc domain to the C-terminus of the second Fc domain (which has three glycine residues added to its N-terminus).

In certain embodiments, a homodimer is provided comprising two such fusion proteins linked via an interchain disulfide bond between their respective GDF15 regions.

More particularly, in a specific embodiment, the homodimer comprises:

(a) two second Fc domains (one each monomer) comprising the sequence (part of the hinge region in parentheses):

GGG (ERKSSVECPCP) APPVAGPSVFLFPPKPKDTLMISRTPEVTCVVV
 DVSHEDPEVQFNWYVDGVEVHNAKTKPREEQFNSTFRVVSVLTVVHQDWL
 NGKEYKCKVSNKGLPAPIEKTISKTKGQPREPQVYTLPPSREEMTKNQVS
 LTCLVKGFYPSDIAVEWESNGQPENNYKTTPMLDSDGSFFLYSKLTVDK
 5 SRWQQGNVFSCSVMHEALHNHYTQKSLSLSPG (SEQ ID NO:28),

(b) two first Fc domains (one each monomer) comprising the sequence (part of the hinge region in parentheses):

(ERKSSVECPCP) APPVAGPSVFLFPPKPKDTLMISRTPEVTCVVVDVS
 HEDPEVQFNWYVDGVEVHNAKTKPREEQFNSTFRVVSVLTVVHQDWLNGK
 10 EYKCKVSNKGLPAPIEKTISKTKGQPREPQVYTLPPSREEMTKNQVSLTC
 LVKGFYPSDIAVEWESNGQPENNYKTTPMLDSDGSFFLYSKLTVDKSRW
 QQGNVFSCSVMHEALHNHYTQKSLSLSPG (SEQ ID NO:29) .

(c) two second polypeptide linkers (one each monomer) comprising the sequence of
 SEQ ID NO:18 linking the N-terminus of the first Fc domain to the C-terminus of the second
 15 Fc domain,

(d) two GDF15 polypeptides (one each monomer) comprising the sequence of SEQ
 ID NO:12, and

(e) two first polypeptide linkers (one each monomer) comprising sequence:

SGGGGSGGGGSGGGGSGGGGS (SEQ ID NO:30)
 20 linking the N-terminus of a GDF15 polypeptide to the C-terminus of a first Fc domain.

In a preferred embodiment, the fusion protein comprises the amino acid sequence
 (part of the hinge region in parentheses; linker sequences double underlined):

GGG (ERKSSVECPCP) APPVAGPSVFLFPPKPKDTLMISRTPEVTCVVV
 DVSHEDPEVQFNWYVDGVEVHNAKTKPREEQFNSTFRVVSVLTVVHQDWL
 25 NGKEYKCKVSNKGLPAPIEKTISKTKGQPREPQVYTLPPSREEMTKNQVS
 LTCLVKGFYPSDIAVEWESNGQPENNYKTTPMLDSDGSFFLYSKLTVDK
 SRWQQGNVFSCSVMHEALHNHYTQKSLSLSPGSGGGGSGGGGSGGGGSGGG
GS (ERKSSVECPCP) APPVAGPSVFLFPPKPKDTLMISRTPEVTCVVVD
 VSHEDPEVQFNWYVDGVEVHNAKTKPREEQFNSTFRVVSVLTVVHQDWLN
 30 GKEYKCKVSNKGLPAPIEKTISKTKGQPREPQVYTLPPSREEMTKNQVSL
 TCLVKGFYPSDIAVEWESNGQPENNYKTTPMLDSDGSFFLYSKLTVDKS
 RWQQGNVFSCSVMHEALHNHYTQKSLSLSPGSGGGGSGGGGSGGGGSGGG
GSARNGDHCPLGPGRCCRLHTVRASLEDLGWADWVLSPREVQVTMCIGAC

PSQFRAANMHAQIKTSLHRLKPDTVPAPCCVPASYNPMVLIQKTDGTVSL
QTYDDLLAKDCHCI (SEQ ID NO:32),

which is encoded by the nucleic acid sequence:

ggaggtggagagcgcaaattcttctgtcgagtgccaccgtgccagcacc
5 acctgtggcaggaccgtcagtccttctcttccccccaaaaccaaggaca
ccctcatgatctcccgaccctgaggtcacgtgctgggtggacgtg
agccacgaagaccccgaggtccagttcaactggtacgtggacggcgtgga
ggtgcataatgccaagacaaaaccacgggaggagcagttcaacagcacgt
tccgtgtggtcagcgtcctcaccgttgtgcaccaggactggctgaacggc
10 aaggagtacaagtgaaggtctccaacaaaggcctcccagccccatcga
gaaaaccatctccaaaaccaaagggcagccccgagaaccacaggtgtaca
ccctgcccccatcccgaggagatgaccaagaaccaggtcagcctgacc
tgcttggtcaaaggttctacccagcgacatcgccgtggagtgggagag
caatgggcagccggagaacaactacaagaccacacctcccatgctggact
15 ccgacggctccttcttctctacagcaagctcaccgtggacaagagcagg
tggcagcaggggaacgtcttctcatgctccgtgatgcatgaggctctgca
caaccactacacgcagaagagcctctccctgtctccgggtggaggtggcg
gtagcgggtggcggaggttcaggtggcgggcgaagcgggtggaggaggttca
gagcggaaatccagcgttgaatgtcctccgtgccctgctccaccctgcgc
20 ggggcctagtgtcttcttttccctccaaaaccaaaggatacactgatga
tcagccggacccccgaggttacgtgcgtcgctcgatgtctcccacgag
gatccagaggtccaattcaactggtacgtggacggggtcgaggtgcataa
tgcaaagacaaagccacgggaagagcagtttaactctactttccgcgtgg
tttctgtgctgaccgtggtgcaccaagattggctcaacggcaaggagtac
25 aagtgaaggtaagcaataaggggtccctgccccattgagaagactat
ctccaagacaaagggacagccacgcgagccacaagtctatacactcccc
cttcccgcgaagaaatgaccaagaatcaggttagcctgacatgcttggtt
aagggtttctacccctctgacatagccgtggagtgggagagcaatggaca
accagagaacaactacaagaccacccacccatgctggatagcgacggtt
30 cattctttctgtatagtaagcttacgtggacaagtcccgggtggcaacaa
ggaaatgtcttttcatgctctgtgatgcacgaggccttgcataatcacta
tactcagaagagcttgagcctcagccccggatctggaggtggcggatccg
ggggcggtggaagcggaggtggtggatcgggaggcgggtggaagcgcgcgc
aacggcgaccactgtccgctcgggcccggacgttgctgccgtctgcacac
35 ggtccgcgcgtcgctggaagacctgggctgggccgattgggtgctgtcgc
cacgggaggtgcaagtgaccatgtgcatcggcgcgtgccccgagccagttc

cgggcggaacatgcacgcgcagatcaagacgagcctgcaccgcctgaa
 gcccgcacgggtgccagcgccctgctgcggtgcccgccagctacaatccca
 tgggtgctcattcaaaagaccgacaccggggtgtcgctccagacctatgat
 gacttggttagccaaagactgccactgcatatga (SEQ ID NO:31).

- 5 As discussed above, in a specific embodiment, a homodimer is provided comprising two monomers having the sequence of SEQ ID NO:32.

II.C. DhHemiFc

- The designations “DhHemi-” or “DhHemiFc domain” in the instant disclosure refer to a polypeptide chain comprising a first Fc domain from which all or part of the hinge region has been removed (“DhFc” domain) linked, directly or via a polypeptide linker, to a second DhFc domain. In particular embodiments, the N-terminal 12 amino acids in the hinge region are removed, *e.g.*, ERKSSVECPCP (SEQ ID NO:15). In one embodiment, the first DhFc domain and the second DhFc domain have the same sequence. In another embodiment, the first DhFc domain and the second DhFc domain have different sequences.

- A fusion protein is provided comprising a DhHemiFc domain and a GDF15 region. Typically, the N-terminus of the GDF15 region is linked, directly or via a polypeptide linker, to the C-terminus of the DhHemiFc domain. However, in some embodiments, the N-terminus of the DhHemiFc domain is linked, directly or via a polypeptide linker, to the C-terminus of the GDF15 region.

In certain embodiments, a dimer is provided comprising two such fusion proteins linked via an interchain disulfide bond between their respective GDF15 regions. In some embodiments the dimer is a homodimer. In other embodiments, the dimer is a heterodimer.

25 *GGGDhFc-(G₄S)₅-DhFc-S(G₄S)₄-GDF15*

- The designation “GGGDhFc-(G₄S)₅-DhFc-S(G₄S)₄-GDF15” in the instant disclosure refers to a fusion protein comprising a GDF15 polypeptide linked to a DhHemiFc domain via a first polypeptide linker comprising the sequence of SEQ ID NO:30, that connects the N-terminus of the GDF15 polypeptide to the C-terminus of the HemiFc domain. The HemiFc domain comprises a first DhFc domain linked to a second DhFc domain via a polypeptide linker comprising the sequence of SEQ ID NO:34, that connects the N-terminus of the first DhFc domain to the C-terminus of the second DhFc domain (which has three glycine residues added to its N-terminus).

- In certain embodiments, a homodimer is provided comprising two such fusion proteins linked via an interchain disulfide bond between their respective GDF15 regions.

More particularly, in a specific embodiment, the homodimer comprises:

(a) two second DhFc domains (one each monomer) comprising the sequence:

GGGAPPVAGPSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVQFNWY
VDGVEVHNAKTKPREEQFNSTFRVSVLTVVHQDWLNGKEYKCKVSNKGL
PAPIEKTISKTKGQPREPQVYTLPPSREEMTKNQVSLTCLVKGFYPSDIA
5 VEWESNGQPENNYKTTPMLDSDGSFFLYSKLTVDKSRWQQGNVFSCSVM
HEALHNHYTQKSLSLSPG (SEQ ID NO:33),

(b) two first DhFc domains (one each monomer) comprising the sequence:

APPVAGPSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVQFNWYVDG
VEVHNAKTKPREEQFNSTFRVSVLTVVHQDWLNGKEYKCKVSNKGLPAP
10 IEKTISKTKGQPREPQVYTLPPSREEMTKNQVSLTCLVKGFYPSDIAVEW
ESNGQPENNYKTTPMLDSDGSFFLYSKLTVDKSRWQQGNVFSCSVMHEA
LHNHYTQKSLSLSPG (SEQ ID NO:35),

(c) two second polypeptide linkers (one each monomer) comprising the sequence:

GGGGSGGGSGGGSGGGSGGGSGGGGS (SEQ ID NO:34)

15 linking the N-terminus of the first DhFc domain to the C-terminus of the second DhFc domain,

(d) two GDF15 polypeptides (one each monomer) comprising the sequence of SEQ ID NO:12, and

(e) two first polypeptide linkers (one each monomer) comprising the sequence of
20 SEQ ID NO:30 linking the N-terminus of a GDF15 polypeptide to the C-terminus of a first DhFc domain.

In a preferred embodiment, the fusion protein comprises the amino acid sequence (linker sequences double underlined):

GGGAPPVAGPSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVQFNWY
25 VDGVEVHNAKTKPREEQFNSTFRVSVLTVVHQDWLNGKEYKCKVSNKGL
PAPIEKTISKTKGQPREPQVYTLPPSREEMTKNQVSLTCLVKGFYPSDIA
VEWESNGQPENNYKTTPMLDSDGSFFLYSKLTVDKSRWQQGNVFSCSVM
HEALHNHYTQKSLSLSPGGGGGSGGGSGGGSGGGSGGGSGGGGSAPPVAGP
SVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVQFNWYVDGVEVHNAK
30 TKPREEQFNSTFRVSVLTVVHQDWLNGKEYKCKVSNKGLPAPIEKTISK
TKGQPREPQVYTLPPSREEMTKNQVSLTCLVKGFYPSDIAVEWESNGQPE
NNYKTTPMLDSDGSFFLYSKLTVDKSRWQQGNVFSCSVMHEALHNHYTQ
KSLSLSPGGGGGSGGGSGGGSGGGSGGGGSARNGDHCPLGPGRCCRLHTR

ASLEDLGWADWVLSPREVQVTMCIGACPSQFRAANMHAQIKTSLHRLKPD
 TVPAPCCVPASYNPMVLIQKTDGTGVSLLQTYDDLLAKDCHCI (SEQ ID NO:37),

which is encoded by the nucleic acid sequence:

```

ggcgggtggagctccgccggtggctggaccctcagtgttcctctttccacc
5 gaagccgaaggacacccttatgattagccggaccccagaggtcacttgcg
tcgtcgtggacgtgtcccatgaggatcccgaagtgcagtttaactggtat
gtggacggagtgagggtccataacgccaaagaccaagccaaggaagaaca
gttcaatagcaccttcgggtggtgtccgtgctcacctggtgcatcaag
actggtgtaatggcaaagagtacaaatgtaagggtgtcaaacaaggggtc
10 ccagcccctattgaaaagaccatctcaaagactaaggacagccacgcga
acctcaagtgtataccctccgccttcacgcgaagaaatgactaagaatc
aggtcagccttacttgtctggtcaagggcttctacccgagcgacattgca
gtcgaatgggagagcaatgggtcagccagagaataactacaagaccactcc
tcccatgcttgatagcgatggaagctttttcctttacagcaagcttactg
15 tggataagtctcgctggcaacagggaaatgtgttcagctgttcagtgatg
catgaagcactccacaatcattacaccagaagtcactcagcctctcacc
cggaggaggaggcgggttctggtggaggagggtctggaggtggagggagcg
gcgagggcgggtctggcgggtggtgggtctgagaggaagtcatcagtggaa
tgcccaccatgccctgctcctcccggtggccggtccgagcgtgtttctctt
20 cccacctaaagcccaaggacactctgatgatctcacggactccggaagtga
cttgtgtggtggtggacgtgtctcatgaggaccctgaagtgcagttcaac
tggtagctggacggcgtggaggtgcacaatgctaagaccaagcctagaga
ggaacagttcaattccacctttcgcgtggtgagcgtcctgaccgtcgtgc
accaggactggcttaacggaaggaatacaagtgaaggtgtccaacaaa
25 ggccttccagctcccattgagaaaaccatctctaaaactaagggtcaacc
aagggaaacccaagtctacaccctccctccgtctagagaagagatgacca
aaaaccagggtgtccctgacctgtctggtgaagggattttaccctcagac
atcgccgtggagtgggaaagcaacggacagccccgaaaacaactataagac
taccctcctatgctggactcagacggatctttcttctctatagcaagc
30 tcaactgtggacaaaatccagatggcaacaagggaatgtgttctcatgcagc
gtgatgcacgaggctcttcacaaccactataaccagaagagcctgtctct
ttcacctggttccggagggtggtgggagcggagggggtggatcaggtggtg
gaggggtccggaggcggaggatccgcacggaatggcgaccactgtccactg
ggacccggaagatgttgtcgccctccacaccgtgagggcctctctggagga
35 ccttggtggtggccgactgggtcctgtcacctcgggaggtccaagtcacca
tgtgtatcggagcctgccccagccaattcagagcagcaaatatgcacgca

```

cagattaagaccagcctgcatcggttaaacctgatactgtgccggctcc
 ttgttgcggtgccagcatcttacaacccgatgggtgctgatccagaaaaccg
 ataccgggtgtctccctccagacttacgacgacctccttgcaaaggactgc
 cattgcatc (SEQ ID NO:36).

- 5 As discussed above, in a specific embodiment, a homodimer is provided comprising two monomers comprising the sequence of SEQ ID NO:37.

II.D. Knob/Hole

- 10 The designations “knob-” or “knobFc domain” in the instant disclosure refer to an Fc domain comprising a “knob” mutation. The designations “hole-” or “holeFc domain” in the instant disclosure refers to a native Fc or Fc variant comprising a “hole” mutation.

- “Knobs” may be created by replacing small amino acid side chains with larger ones and “holes” may be created by replacing large amino acid side chains with smaller ones. *See, e.g.,* Ridgway JBB 1996, *Protein Eng.* 9:617-621; Merchant AM 1998, *Nature Biotech.*
 15 16:677-681.

- In one embodiment, a “knobFc” domain is provided by introducing a threonine to tryptophan mutation (T366W) in the sequence of an Fc domain. In one embodiment, a “holeFc” domain is provided by introducing a threonine to serine mutation (T366S), a leucine to alanine mutation (L368A) and a tyrosine to valine mutation (Y407V) in the sequence of an
 20 Fc domain.

The C-terminal lysine (K447) optionally may be deleted in the knobFc domain, the holeFc domain, or both. This may be advantageous, for example, when a peptide is fused at the C-terminus to reduce proteolysis of the fusion protein.

- In one embodiment, a “DhknobFc” domain is provided by introducing a threonine to
 25 tryptophan mutation (T366W) in the sequence of an Fc domain from which all or part of the hinge region has been removed. In one embodiment, a “DhholeFc” domain is provided by introducing a threonine to serine mutation (T366S), a leucine to alanine mutation (L368A) and a tyrosine to valine mutation (Y407V) in the sequence of an Fc domain from which all or part of the hinge region has been removed.

- 30 The C-terminal lysine (K447) optionally may be deleted in the DhknobFc, the DhholeFc, or both. This may be advantageous, for example, when a peptide is fused at the C-terminus to reduce proteolysis of the fusion protein.

- In some embodiments, a heterodimer is providing comprising (i) a first polypeptide chain comprising a GDF15 region linked to a holeFc domain directly or via a polypeptide
 35 linker and (ii) a second polypeptide chain comprising a knobFc domain. In other embodiments, a heterodimer is provided comprising (i) a first polypeptide chain comprising a

GDF15 region linked to a knobFc domain, directly or via a polypeptide linker and (ii) a second polypeptide chain comprising a holeFc domain.

In some embodiments, a heterodimer is provided comprising (i) a first polypeptide chain comprising a GDF15 region linked to a DhholeFc domain directly or via a polypeptide linker and (ii) a second polypeptide chain comprising a DhknobFc domain. In other embodiments, a heterodimer is provided comprising (i) a first polypeptide chain comprising a GDF15 region linked to a DhknobFc domain, directly or via a polypeptide linker and (ii) a second polypeptide chain comprising a DhholeFc domain.

In some embodiments, a tetramer is provided comprising two such heterodimers in which the heterodimers are linked via an interchain disulfide bond between the GDF15 regions of their respective first polypeptide chains. See Figure 1 for a graphic depiction of an embodiment of a heterotetramer comprising two heterodimers, where each heterodimer comprises (i) a first polypeptide chain comprising a GDF15 polypeptide linked to a DhknobFc domain via a polypeptide linker and (ii) a second polypeptide chain comprising a DhholeFc domain.

DhknobFc-(G₄S)₄-GDF15:DhholeFc

The designation “DhknobFc-(G₄S)₄-GDF15:DhholeFc” in the instant disclosure refers to a heterodimer comprising (i) a first polypeptide chain comprising a GDF15 polypeptide linked to a DhknobFc domain via a linker comprising the sequence of SEQ ID NO:18 that connects the N-terminus of the GDF15 polypeptide to the C-terminus of the DhknobFc domain and (ii) a polypeptide chain comprising a DhholeFc domain.

In certain embodiments, a tetramer is provided comprising a dimer of two DhknobFc-(G₄S)₄-GDF15:DhholeFc heterodimers in which the first polypeptide chains of each heterodimer are linked via an interchain disulfide bond between their respective GDF15 regions.

More particularly, in a specific embodiment, the tetramer comprises:

(a) two DhknobFc domains (one each heterodimer) comprising the sequence:

APELLGGPSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVD
GVEVHNAKTKPREEQYNSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPA
PIEKTISKAKGQPREPQVYTLPPSRDELTKNQVSLWCLVKGFYPSDIAVE
WESNGQPENNYKTTTPVLDSDGSFFLYSKLTVDKSRWQQGNVFSQSVME
ALHNHYTQKSLSLSPG (SEQ ID NO:16),

(b) two DhholeFc domains (one each heterodimer) comprising the sequence:

APELLGGPSVFLFPPKPKDTLMISRTPEVTCVVDVSHEDPEVKFNWYVD
 GVEVHNAKTKPREEQYNSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPA
 PIEKTISKAKGQPREPQVYTLPPSRDELTKNQVSLSCAVKGFYPSDIAVE
 WESNGQPENNYKTTPPVLDSDGSFFLVSKLTVDKSRWQQGNVFSQVMHE

5 ALHNHYTQKSLSLSPG (SEQ ID NO:281),

(c) two GDF15 polypeptides (one each heterodimer) comprising the sequence of SEQ ID NO:12, and

(d) two polypeptide linkers (one each heterodimer) comprising the sequence of SEQ ID NO:18 each linking the N-terminus of a GDF15 polypeptide to the C-terminus of a
 10 DhknobFc domain via peptide bonds.

In a preferred embodiment, the first polypeptide chain comprises the amino acid sequence (linker sequence double underlined):

APELLGGPSVFLFPPKPKDTLMISRTPEVTCVVDVSHEDPEVKFNWYVD
 GVEVHNAKTKPREEQYNSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPA
 15 PIEKTISKAKGQPREPQVYTLPPSRDELTKNQVSLWCLVKGFYPSDIAVE
 WESNGQPENNYKTTPPVLDSDGSFFLYSKLTVDKSRWQQGNVFSQVMHE
 ALHNHYTQKSLSLSPGGGGGGSGGGGSGGGGSGGGGSARNGDHCPLGPGRC
 CRLHTVRASLEDLGWADWVLSPREVQVTCIGACPSQFRAANMHAQIKTS
 LHLKPDTPVAPCCVPASYNPMVLIQKTDGTGVSQTQYDDLLAKDCHCI (SEQ ID
 20 NO:20),

which is encoded by the nucleic acid sequence:

gcacctgaactcctggggggaccgtcagtccttcctcttccccccaaaacc
 caaggacaccctcatgatctcccgaccctgaggtcacatgcgtggtgg
 tggacgtgagccacgaagaccctgaggtcaagttcaactggtacgtggac
 25 ggcgtggaggtgcataatgccaaagacaaagccgcgggaggagcagtacaa
 cagcacgtaccgtgtggtcagcgtcctcaccgtcctgcaccaggactggc
 tgaatggcaaggagtacaagtgaaggtctccaacaaagccctcccagcc
 cccatcgagaaaaccatctccaaagccaaagggcagccccgagaaccaca
 ggtgtacaccctgcccccatcccggtatgagctgaccaagaaccagggtca
 30 gcctgtggtgcctggtcaaaggcttctatcccagcgacatcgccgtggag
 tgggagagcaatgggcagccggagaacaactacaagaccacgcctcccgt
 gctggactccgacggctccttcttctctacagcaagctcaccgtggaca
 agagcaggtggcagcaggggaacgtcttctcatgctccgtgatgcatgag
 gctctgcacaaccactacacgcagaagagcctctccctgtctccgggtgg
 35 aggtggtggatccggaggcgggtggaagcggaggtggtggatctggaggcg

gtggaagcgcgcgcaacggagaccactgtccgctcgggccccgggcgttgc
 tgccgtctgcacacgggtccgcgcgtcgctggaagacctgggctgggcga
 ttgggtgctgtcgccacgggaggtgcaagtgacctgtgcatcggcgct
 gcccagaccagttccgggcggaacatgcacgcgcagatcaagacgagc
 5 ctgcaccgcctgaagcccgacacgggtgccagcgccctgctgctgcccgc
 cagctacaatcccatggtgctcattcaaaagaccgacaccggggtgtcgc
 tccagacctatgatgacttgttagccaaagactgccactgcatatga (SEQ ID
 NO:19) .

In a preferred embodiment, the second polypeptide chain comprises the amino acid sequence:

10 APELLGGPSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVD
 GVEVHNAKTKPREEQYNSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPA
 PIEKTISKAKGQPREPQVYTLPPSRDELTKNQVSLSCAVKGFYPSDIAVE
 WESNGQPENNYKTTTPVLDSDGSFFLVSKLTVDKSRWQQGNVFSQVMHE
 ALHNHYTQKSLSLSPGK (SEQ ID NO:17) ,

15 which is encoded by the nucleic acid sequence:

gcacctgaactcctggggggaccgtcagtccttctcttccccccaaaacc
 caaggacaccctcatgatctcccgaccctgaggtcacatgcgtggtgg
 tggacgtgagccacgaagaccctgaggtcaagttcaactggtacgtggac
 ggctgagggtgcataatgccaaagacaaagccgcgggaggagcagtacaa
 20 cagcacgtaccgtgtggtcagcgtcctcaccgtcctgcaccaggactggc
 tgaatggcaaggagtacaagtgcaaggtctccaacaaagccctcccagcc
 cccatcgagaaaaccatctccaaagccaaagggcagccccgagaaccaca
 ggtgtacaccctgcccccatcccggtatgagctgaccaagaaccaggtca
 gcctgagctgcgcggtcaaaggcttctatcccagcgacatcgccgtggag
 25 tgggagagcaatgggcagccggagaacaactacaagaccacgcctcccg
 gctggactccgacggctccttcttctcgtcagcaagctcaccgtggaca
 agagcaggtggcagcaggggaacgtcttctcatgctccgtgatgcatgag
 gctctgcacaaccactacacgcagaagagcctctccctgtctccgggtaa
 atga (SEQ ID NO:21)

30 As discussed above, in a specific embodiment, a tetramer is provided comprising two monomers having the sequence of SEQ ID NO:20 and two monomers having the sequence of SEQ ID NO:17.

II.E. Charged pair (delHinge)

The designation "CpmFc(+)" domain" in the instant disclosure refers to an Fc domain comprising a "positive" charged pair mutation. The designation "CpmFc(-)" domain" in the instant disclosure refers to an Fc domain comprising a "negative" charged pair mutation. Note that use of the terms "positive" and "negative" is for ease of reference (i.e., to describe the nature of the charged pair mutations in the Fc domains) and not to indicate that the overall sequence or construct necessarily has a positive or negative charge.

In one embodiment, a "positive" charged pair mutation is provided by introducing a glutamic acid to lysine mutation (E356K) and an aspartic acid to lysine mutation (D399K) in the sequence of an Fc domain. In one embodiment, a "negative" charged pair mutation is provided by introducing two lysine to aspartic acid mutations (K392D, K409D) in the sequence of an Fc domain.

When incubated together, the aspartate residues associate with the lysine residues through electrostatic force, facilitating formation of heterodimers between the CpmFc(+) domains and CpmFc(-) domains, and reducing or preventing formation of homodimers between the CpmFc(+) sequences or between CpmFc(-) sequences.

The C-terminal lysine (K447) optionally may be deleted in the CpmFc(+) domain, the CpmFc(-) domain, or both. This may be advantageous, for example, when a peptide is fused at the C-terminus to reduce proteolysis of the fusion protein.

In one embodiment, a "DhCpmFc(+)" domain is provided by introducing a glutamic acid to lysine mutation (E356K) and an aspartic acid to lysine mutation (D399K) in the sequence of an Fc domain from which all or part of the hinge region has been removed. In one embodiment, a "DhCpmFc(-)" domain is provided by introducing two lysine to aspartic acid mutations (K392D, K409D) in the sequence of an Fc domain from which all or part of the hinge region has been removed.

When incubated together, the aspartate residues associate with the lysine residues through electrostatic force, facilitating formation of heterodimers between the DhCpmFc(+) domains and DhCpmFc(-) domains, and reducing or preventing formation of homodimers between the DhCpmFc(+) sequences or between DhCpmFc(-) sequences.

The C-terminal lysine (K447) optionally may be deleted in the DhCpmFc(+), the DhCpmFc(-), or both. This may be advantageous, for example, when a peptide is fused at the C-terminus to reduce proteolysis of the fusion protein.

In some embodiments, a heterodimer is provided comprising (i) a first polypeptide chain comprising a GDF15 region linked to a CpmFc(+) domain, and (ii) a second polypeptide chain comprising a CpmFc(-) domain. In some embodiments, the N-terminus of the GDF15 region is linked to the C-terminus of the CpmFc(+) domain, directly or via a polypeptide linker. In other embodiments, the N-terminus of the CpmFc(+) domain is linked to the C-terminus of the GDF15 region, directly or via a polypeptide linker.

In other embodiments, a heterodimer is provided comprising (i) a first polypeptide chain comprising a GDF15 region linked to a CpmFc(-) domain, and (ii) a second polypeptide chain comprising a CpmFc(+) domain. In some embodiments, the N-terminus of the GDF15 region is linked to the C-terminus of the CpmFc(-) domain, directly or via a polypeptide linker. In other embodiments, the N-terminus of the CpmFc(-) domain is linked to the C-terminus of the GDF15 region, directly or via a polypeptide linker.

In some embodiments, a heterodimer is provided comprising (i) a first polypeptide chain comprising a GDF15 region linked to a DhCpmFc(+) domain, and (ii) a second polypeptide chain comprising a DhCpmFc(-) domain. In some embodiments, the N-terminus of the GDF15 region is linked to the C-terminus of the DhCpmFc(+) domain, directly or via a polypeptide linker. In other embodiments, the N-terminus of the DhCpmFc(+) domain is linked to the C-terminus of the GDF15 region, directly or via a polypeptide linker.

In other embodiments, a heterodimer is provided comprising (i) a first polypeptide chain comprising a GDF15 region linked to a DhCpmFc(-) domain, and (ii) a second polypeptide chain comprising a DhCpmFc(+) domain. In some embodiments, the N-terminus of the GDF15 region is linked to the C-terminus of the DhCpmFc(-) domain, directly or via a polypeptide linker. In other embodiments, the N-terminus of the DhCpmFc(-) domain is linked to the C-terminus of the GDF15 region, directly or via a polypeptide linker.

In some embodiments, a tetramer is provided comprising a dimer of two such heterodimers in which the two first polypeptide chains of the heterodimers are linked via an interchain disulfide bond between their respective GDF15 regions. See Figure 4 for a graphic depiction of an embodiment of a tetramer comprising two heterodimers, where each heterodimer comprises (i) a first polypeptide chain comprising a GDF15 polypeptide linked via a polypeptide linker to a DhCpmFc(-) domain and (ii) a second polypeptide chain comprising a DhCpmFc(+) domain.

II.E.1 DhCpmFc(+)-(1K)-GDF15:DhCpmFc(-)

The designation “DhCpmFc(+)-(1K)-GDF15:DhCpmFc(-)” in the instant disclosure refers to a heterodimer comprising (i) a first polypeptide chain comprising a GDF15 polypeptide linked to a DhCpmFc(+) domain via a linker comprising the sequence of SEQ ID NO:40 that connects the N-terminus of the GDF15 polypeptide to the C-terminus of the DhCpmFc(+) domain and (ii) a second polypeptide chain comprising a DhCpmFc(-) domain.

In certain embodiments, a tetramer is provided comprising a dimer of two DhCpmFc(+)-(1K)-GDF15:DhCpmFc(-) heterodimers in which the two first polypeptide chains of each heterodimer are linked via an interchain disulfide bond between their respective GDF15 regions.

More particularly, in a specific embodiment, the tetramer comprises:

(a) two DhCpmFc(+) domains (one each heterodimer) comprising the sequence:

APELLGGPSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVD
 GVEVHNAKTKPREEQYNSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPA
 PIEKTISKAKGQPREPQVYTLPPSRKEMTKNQVSLTCLVKGFYPSDIAVE
 5 WESNGQPENNYKTTPPVLKSDGSFFLYSKLTVDKSRWQQGNVFSQVMHE
 ALHNHYTQKSLSLSPG (SEQ ID NO:38),

(b) two DhCpmFc(-) domains (one each heterodimer) comprising the sequence:

APELLGGPSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVD
 GVEVHNAKTKPREEQYNSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPA
 10 PIEKTISKAKGQPREPQVYTLPPSREEMTKNQVSLTCLVKGFYPSDIAVE
 WESNGQPENNYDTTPPVLDSDGSFFLYSDLTVDKSRWQQGNVFSQVMHE
 ALHNHYTQKSLSLSPG (SEQ ID NO:282),

(c) two GDF15 polypeptides (one each heterodimer) comprising the sequence of SEQ ID NO:12, and

15 (d) two polypeptide linkers (one each heterodimer) comprising the sequence:

GGSGATGGSGSVASSGSGSATHL (SEQ ID NO:40)

each linking the N-terminus of a GDF15 polypeptide to the C-terminus of a DhCpmFc(+) domain via a peptide bond.

20 In a preferred embodiment, the first polypeptide chain comprises the amino acid sequence (linker sequence double underlined):

APELLGGPSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVD
 GVEVHNAKTKPREEQYNSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPA
 PIEKTISKAKGQPREPQVYTLPPSRKEMTKNQVSLTCLVKGFYPSDIAVE
 WESNGQPENNYKTTPPVLKSDGSFFLYSKLTVDKSRWQQGNVFSQVMHE
 25 ALHNHYTQKSLSLSPGGGSGATGGSGSVASSGSGSATHLLARNGDHCPLGP
 GRCCRLHTVRASLEDLGWADWVLSPREVQVTMCIGACPSQFRAANMHAQI
 KTSLHRLKPDTPAPCCVPASYNPMVLIQKTDGTGVSQTYYDLLAKDCHC
 I (SEQ ID NO:42),

which is encoded by the nucleic acid sequence:

30 gccccagagctgcttggtggaccatccgtgttcctgtttcctccaaagcc
 gaaggacaccctgatgatctcaagaactccggaagtgacttgcgctcgctg
 tggacgtgtcacatgaggatccagaggtcaagttcaattggtatgtggac

ggagtggaagtgcataacgccaagaccaaaccggaagaacagtacaa
 tagcacctaccgctggtgagcgctccttactgtgctccaccaggactggc
 ttaatgggaaggaatacaagtgttaaggtgtccaacaaggccctccccgct
 cccatcgaaaagaccatctcaaaggcaaagggaaccaagggaacctca
 5 agtgtagaccctgcctccgagcaggaaggagatgaccaagaaccaggtea
 gcctgacttgtctcgtgaagggcttctatcccagcgatattgctgtggaa
 tgggagtgaaatggccagccccgagaataactacaaaactacccccaccgt
 gctgaaatctgatgggtccttcttcttcttactccaagctgaccgtggaca
 agagccgctggcaacaaggcaatgtcttttagctgctcagtgatgcatgag
 10 gctctccataatcactacactcagaagtcactgtccctgtcacctggcgg
 atccggttctgctactgggtggttccggctccgtcgcaagctctggttcag
 gcagtgcgactcatctggcacggaacggggaccattgtcccctgggacct
 ggtcgggtgctgccggcttcacaccgtcagagcctctctggaggaccttg
 atgggctgattgggtgctgagccctcgggaggtgcaagtcaccatgtgca
 15 tcggggcctgccctagccagttccgcgcagccaacatgcacgctcagatc
 aaaacctctcttcacagactgaagccccgacaccgtgccagcaccttgctg
 tgtgccggcctcttataaccccatggctcctcattcagaaaaccgacaccg
 gagtgtcacttcagacttacgatgacctcctggccaaggactgccactgc
 ata (SEQ ID NO:41) .

20 In a preferred embodiment, the second polypeptide chain and comprises the amino
 acid sequence

APELLGGPSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVD
 GVEVHNAKTKPREEQYNSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPA
 PIEKTISKAKGQPREPQVYTLPPSREEMTKNQVSLTCLVKGFYPSDIAVE
 25 WESNGQPENNYDTPPVLDSDGSFFLYSDLTVDKSRWQQGNVVFSCSVMHE
 ALHNHYTQKSLSLSPGK (SEQ ID NO:39) ,

which is encoded by the nucleic acid sequence:

gcgccggaactgctgggcggccccgagcgtgtttctgtttccgccgaaacc
 gaaagataccctgatgattagccgcacccccggaagtgcctgcgtggtgg
 30 tggatgtgagccatgaagatccggaagtgaaatttaactggatgtggat
 ggcgtggaagtgcataacgcgaaaaccaaaccgcgcaagaacagtataa
 cagcacctatcgctggtgagcgtgctgaccgtgctgcatcaggattggc
 tgaacggcaaagaatataaatgcaaagtgagcaaaaagcgctgccggcg
 ccgattgaaaaaaccattagcaaagcgaaaggccagccgcgcaaccgca
 35 ggtgtataccctgccgccgagccgcgaagaaatgacaaaaaccagggtga

gcctgacctgcctggtgaaaggcttttatccgagcgatattgcggtggaa
 tgggaaagcaacggccagccggaacaactatgataccaccccgccggt
 gctggatagcgatggcagcttttttctgtatagcgatctgaccgtggata
 aaagccgctggcagcagggcaacgtgttttagctgcagcgtgatgcatgaa
 5 gcgctgcataaccattatacccagaaaaagcctgagcctgagcccgggcaa
 a (SEQ ID NO:43).

As discussed above, a tetramer is provided comprising two polypeptide chains comprising the sequence of SEQ ID NO:42 and two polypeptide chains comprising the sequence of SEQ ID NO:39.

10

II.E.2 DhCpmFc(-)-GDF15:DhCpmFc(+)

The designation “DhCpmFc(-)-GDF15:DhCpmFc(+)” in the instant disclosure refers to a heterodimer comprising (i) a first polypeptide chain comprising a GDF15 polypeptide, the N-terminus of which is linked directly to the C-terminus of a DhCpmFc(-) domain, and
 15 (ii) a second polypeptide chain comprising a DhCpmFc(+) domain.

In certain embodiments, a tetramer is provided comprising a dimer of two DhCpmFc(-)-GDF15:DhCpmFc(+) heterodimers in which the two first polypeptide chains of each heterodimer are linked via an interchain disulfide bond between their respective GDF15 regions.

20

More particularly, in a specific embodiment, the tetramer comprises:

(a) two DhCpmFc(+) domains (one each heterodimer) comprising the amino acid sequence:

APELLGGPSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVD
 GVEVHNAKTKPREEQYNSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPA
 25 PIEKTISKAKGQPREPQVYTLPPSRKEMTKNQVSLTCLVKGFYPSDIAVE
 WESNGQPENNYKTTTPVLKSDGSFFLYSKLTVDKSRWQQGNVFSQVMHE
 ALHNHYTQKSLSLSPG (SEQ ID NO:283),

(b) two DhCpmFc(-) domains (one each heterodimer) comprising the amino acid sequence:

APELLGGPSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVD
 GVEVHNAKTKPREEQYNSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPA
 30 PIEKTISKAKGQPREPQVYTLPPSREEMTKNQVSLTCLVKGFYPSDIAVE
 WESNGQPENNYDTTPVLDSDGSFFLYSDLTVDKSRWQQGNVFSQVMHE
 ALHNHYTQKSLSLSPG (SEQ ID NO:48),

and,

(c) two GDF15 polypeptides (one each heterodimer) comprising the sequence of SEQ ID NO:12.

In a preferred embodiment, the first polypeptide chain comprises the amino acid
5 sequence:

APELLGGPSVFLFPPKPKDTLMI SRTPEVTCVVVDVSHEDPEVKFNWYVD
GVEVHNAKTKPREEQYNSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPA
PIEKTISKAKGQPREPQVYTLPPSREEMTKNQVSLTCLVKGFYPSDIAVE
WESNGQPENNYDTPPVLDSDGSFFLYSDLTVDKSRWQQGNVFCFSVMHE
10 ALHNHYTQKSLSLSPGARNGDHCPLGPGRCCRLHTVRASLEDLGWADWVL
SPREVQVMTMCIGACPSQFRAANMHAQIKTSLHRLKPDTPAPCCVPASYN
PMVLIQKTDGTGVSLSQTYDDLAKDCHCI (SEQ ID NO:50),

which is encoded by the nucleic acid sequence:

gcacctgaactcctggggggaccgtcagtccttcctcttccccccaaaacc
15 caaggacaccctcatgatctcccgaccctgaggtcacatgcgtggtgg
tggacgtgagccacgaagaccctgaggtcaagttcaactggtacgtggac
ggcgtggaggtgcataatgccaaagacaaagccgcgggaggagcagtacaa
cagcacgtaccgtgtggtcagcgtcctcaccgtcctgcaccaggactggc
tgaatggcaaggagtacaagtgaaggtctccaacaagccctcccagcc
20 cccatcgagaaaaccatctccaaagccaaagggcagccccgagaaccaca
ggtgtacaccctgcccccatcccgaggagatgaccaagaaccagggtca
gcctgacctgcctggtcaaaggcttctatcccagcgacatcgccgtggag
tgggagagcaatgggcagccggagaacaactacgacaccacgcctcccgt
gctggactccgacggctccttcttctctatagcgacctcaccgtggaca
25 agagcaggtggcagcaggggaacgtcttctcatgctccgtgatgcatgag
gctctgcacaaccactacacgcagaagagcctctccctgtctccgggtgc
gcgcaacggagaccactgtccgctcgggcccggcggttgctgccgtctgc
acacggtccgcgcgtcgctggaagacctgggctgggccgattgggtgctg
tcgccacgggaggtgcaagtgacatgtgcatcggcgcgtgcccgagcca
30 gttccggggcggaacatgcacgcgcagatcaagacgagcctgcaccgcc
tgaagcccgacacggtgccagcgcctgctgcgtgcccgcagctacaat
cccatggtgctcattcaaaagaccgacaccggggtgtcgctccagacct
tgatgacttgtagccaaagactgccactgcata (SEQ ID NO:49).

In a preferred embodiment, the second polypeptide chain comprises the amino acid
35 sequence:

APELLGGPSVFLFPPKPKDTLMISRTPEVTCVVDVSHEDPEVKFNWYVD
 GVEVHNAKTKPREEQYNSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPA
 PIEKTIKAKGQPREPQVYTLPPSRKEMTKNQVSLTCLVKGFYPSDIAVE
 WESNGQPENNYKTTPPVLKSDGSAFLYSLKLTVDKSRWQQGNVVFSCVMHE
 5 ALHNHYTQKSLSLSPGK (SEQ ID NO:47),

which is encoded by the nucleic acid sequence:

gcacctgaactcctggggggaccgtcagtccttcctcttccccccaaaacc
 caaggacaccctcatgatctcccggaacctgaggtcacatgcgtggtgg
 tggacgtgagccacgaagaccctgaggtcaagttcaactggtacgtggac
 10 ggcgtggaggtgcataatgccaaagacaaagccgcggaggagcagtacaa
 cagcacgtaccgtgtggtcagcgtcctcaccgtcctgcaccaggactggc
 tgaatggcaaggagtacaagtgaaggtctccaacaaagccctcccagcc
 cccatcgagaaaaccatctccaaagccaaagggcagccccgagaaccaca
 ggtgtacaccctgcccccatcccgaaggagatgaccaagaaccagggtca
 15 gcctgacctgcctggtcaaaggcttctatcccagcgacatcgccgtggag
 tgggagagcaatgggcagccggagaacaactacaagaccacgcctcccgt
 gctgaagtccgacggctccttcttctctatagcaagctcaccgtggaca
 agagcaggtggcagcaggggaacgtcttctcatgctccgtgatgcatgag
 gctctgcacaaccactacacgcagaagagcctctccctgtctccgggtaa
 20 atga (SEQ ID NO:51).

As discussed above, a tetramer is provided comprising two polypeptide chains having the sequence of SEQ ID NO:50 and two polypeptide chains having the sequence of SEQ ID NO:47.

25 II.E.3 *DhCpmFc(-)-GDF15(N3D):DhCpmFc(+)*

The designation “*DhCpmFc(-)-GDF15(N3D):DhCpmFc(+)*” in the instant disclosure refers to a heterodimer comprising (i) a first polypeptide chain comprising a naturally-occurring variant of a GDF15 having an asparagine to aspartic acid mutation (N3D) (“GDF15(N3D)”), the N-terminus of which is linked directly to the C-terminus of a
 30 *DhCpmFc(-)* domain, and (ii) a second polypeptide chain comprising a *DhCpmFc(+)* domain. The N3D mutation may reduce deamidation induced heterogeneity.

In certain embodiments, a tetramer is provided comprising a dimer of two *DhCpmFc(-)-GDF15(N3D):DhCpmFc(+)* heterodimers in which the two first polypeptide chains of each heterodimer are linked via an interchain disulfide bond between their
 35 respective GDF15 regions.

More particularly, in a specific embodiment, the tetramer comprises:

(a) two DhCpmFc(+) domains (one each heterodimer) having the sequence of SEQ ID NO:283,

(b) two DhCpmFc(-) domains (one each heterodimer) having the sequence of SEQ ID NO:48, and

(c) two GDF15(N3D) polypeptides (one each heterodimer) comprising the sequence:

ARDGDHCPLGPGRCRLHTVRASLEDLGWADWVLSPREVQVTCIGACPS
QFRAANMHAQIKTSLHRLKPDTVPAPCCVPASYNPMVLIQKTDGTGVSQT
YDDLAKDCHCI (SEQ ID NO:52) .

In a preferred embodiment, the first polypeptide chain comprises the amino acid sequence:

APELLGGPSVFLFPPKPKDTLMI SRTPEVTCVVVDVSHEDPEVKFNWYVD
GVEVHNAKTKPREEQYNSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPA
PIEKTISKAKGQPREPQVYTLPPSREEMTKNQVSLTCLVKGFYPSDIAVE
WESNGQPENNYDTTPPVLDSDGSFFLYSDLTVDKSRWQQGNVFSQSVME
ALHNHYTQKSLSLSPGARDGDHCPLGPGRCRLHTVRASLEDLGWADWVLS
SPREVQVTCIGACPSQFRAANMHAQIKTSLHRLKPDTVPAPCCVPASYN
PMVLIQKTDGTGVSQTYDDLAKDCHCI (SEQ ID NO:54) ,

which is encoded by the nucleic acid sequence:

gcacctgaactcctggggggaccgtcagtccttcctcttccccccaaaacc
caaggacaccctcatgatctcccggaacctgaggtcacatgcgtggtgg
tggacgtgagccacgaagacctgaggtcaagttcaactggtacgtggac
ggcgtggaggtgcataatgccaaagacaaagccgcgggaggagcagtacaa
cagcacgtaccgtgtggtcagcgtcctcaccgtcctgcaccaggactggc
tgaatggcaaggagtacaagtgaaggtctccaacaaagccctcccagcc
cccatcgagaaaaccatctccaaagccaaagggcagccccgagaaccaca
ggtgtacaccctgcccccatcccgaggagatgaccaagaaccaggtca
gcctgacctgcctggtcaaaggcttctatcccagcgacatcgccgtggag
tgggagagcaatgggcagccggagaacaactacgacaccacgcctcccgt
gctggactccgacggctccttcttcctctatagcgacctaccgtggaca
agagcaggtggcagcaggggaacgtcttctcatgctccgtgatgcatgag
gctctgcacaaccactacacgcagaagagcctctccctgtctccgggtgc
gcgcgacggagaccactgtccgctcgggccccggcggttgctgccgtctgc
aacgggtccgcgcgtcgctggaagacctgggctgggcccattgggtgctg

tcgccacgggaggtgcaagtgaccatgtgcatcggcgcgtgcccagagcca
 gttccgggcggaacatgcacgcgcagatcaagacgagcctgcaccgcc
 tgaagccccgacacggtgccagcgccctgctgctgcccgccagctacaat
 cccatggtgctcattcaaaagaccgacaccggggtgtcgtccagaccta
 5 tgatgacttgtagccaaagactgccactgcata (SEQ ID NO:53).

In a preferred embodiment, the second polypeptide chain comprises the amino acid sequence of SEQ ID NO:47, which is encoded by the nucleic acid sequence of SEQ ID NO:51.

As discussed above, a tetramer is provided comprising two polypeptide chains having
 10 the sequence of SEQ ID NO:54 and two polypeptide chains having the sequence of SEQ ID NO:47.

II.E.4 DhCpmFc(-)-GDF15(Ndel3):DhCpmFc(+)

The designation “DhCpmFc(-)-GDF15(Ndel3):DhCpmFc(+)” in the instant
 15 disclosure refers to a heterodimer comprising (i) a first polypeptide chain comprising a variant of GDF15 in which the first three amino acids are deleted (“GDF15(Ndel3)”, the N-terminus of which is linked directly to the C-terminus of a DhCpmFc(-) domain, and (ii) a second polypeptide chain comprising a DhCpmFc(+) domain. The Ndel3 variant may reduce N3 deamidation and subsequent D3 isomerization induced heterogeneity.

20 In certain embodiments, a tetramer is provided, comprising a dimer of two DhCpmFc(-)-GDF15(Ndel3):DhCpmFc(+) heterodimers in which the two first polypeptide chains of each heterodimer are linked via an interchain disulfide bond between their respective GDF15 regions.

More particularly, in a specific embodiment, the tetramer comprises:

25 (a) two DhCpmFc(+) domains (one each heterodimer) having the sequence of SEQ ID NO:283,

(b) two DhCpmFc(-) domains (one each heterodimer) having the sequence of SEQ ID NO:48, and

(c) two GDF15(Ndel3) polypeptides (one each heterodimer) having the sequence:

30 GDHCPLGPGRCRLHTVRASLEDLGWADWVLSPREVQVTMCIGACPSQFR
 AANMHAQIKTSLHRLKPDTPAPCCVPASYNPMVLIQKTDGTGVSLQTYDD
 LLAKDCHCI (SEQ ID NO:55).

In a preferred embodiment, the first polypeptide chain comprises the amino acid sequence:

APELLGGPSVFLFPPKPKDTLMISRTPEVTCVVDVSHEDPEVKFNWYVD
 GVEVHNAKTKPREEQYNSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPA
 PIEKTISKAKGQPREPQVYTLPPSREEMTKNQVSLTCLVKGFYPSDIAVE
 WESNGQPENNYDTPPVLDSDGSFFLYSDLTVDKSRWQQGNVFSQSVNHE
 5 ALHNHYTQKSLSLSPGGDHCPLGPGRCRLHTVRSLEDLGWADWVLSPR
 EVQVTCIGACPSQFRAANMHAQIKTSLHRLKPDTPAPCCVPASYNPMV
 LIQKTDGTGVSLSQTYDDLLAKDCHCI (SEQ ID NO:57),

which is encoded by the nucleic acid sequence:

gcacctgaactcctggggggaccgtcagtccttcctcttccccccaaaacc
 10 caaggacaccctcatgatctcccgaccctgaggtcacatgcgtggtgg
 tggacgtgagccacgaagaccctgaggtcaagttcaactggtacgtggac
 ggcgtggaggtgcataatgccaaagacaaagccgcgggaggagcagtacaa
 cagcacgtaccgtgtggtcagcgtcctcaccgtcctgcaccaggactggc
 tgaatggcaaggagtacaagtgaaggtctccaacaaagccctcccagcc
 15 cccatcgagaaaaccatctccaaagccaaagggcagccccgagaaccaca
 ggtgtacaccctgcccccatcccgaggagatgaccaagaaccagggtca
 gcctgacctgcctggtcaaaggcttctatcccagcgacatcgccgtggag
 tgggagagcaatgggcagccggagaacaactacgacaccacgcctcccg
 gctggactccgacggctccttcttctctatagcgacctcaccgtggaca
 20 agagcaggtggcagcaggggaacgtcttctcatgctccgtgatgcagag
 gctctgcacaaccactacacgcagaagagcctctccctgtctccgggtgg
 agaccactgtccgctcggggcccggttgcctgctgacacgggtcc
 gcgcgtcgctggaagacctgggctgggcccattgggtgctgctgccacgg
 gaggtgcaagtgacctgtgcatcggcgcgtgcccagaccagttccgggc
 25 ggcaaacatgcacgcgcagatcaagacgagcctgcaccgcctgaagccc
 acacggtgccagcgccctgctgctgcccgcagctacaatcccatggtg
 ctcatcctaaaagaccgacaccgggtgtcgtccagacctatgatgactt
 gtttagccaaagactgccactgcata (SEQ ID NO:56).

In a preferred embodiment, the second polypeptide chain comprises the amino acid
 30 sequence of SEQ ID NO:47, which is encoded by the nucleic acid sequence of SEQ ID
 NO:51.

As discussed above, a tetramer is provide comprising two polypeptide chains having
 the sequence of SEQ ID NO:57 and two polypeptide chains having the sequence of SEQ ID
 NO:47.

35

II.E.5 DhCpmFc(-)-G₄-GDF15(N3D):DhCpmFc(+)

The designation “DhCpmFc(-)-G₄-GDF15(N3D):DhCpmFc(+)” in the instant disclosure refers to a heterodimer comprising (i) a first polypeptide chain comprising a GDF15(N3D) polypeptide linked to a DhCpmFc(-) domain via a linker comprising the sequence of SEQ ID NO:58 that connects the N-terminus of the GDF15(N3D) polypeptide to the C-terminus of the DhCpmFc(-) domain and (ii) a second polypeptide chain comprising a DhCpmFc(+) domain.

In certain embodiments, a tetramer is provided, comprising a dimer of two DhCpmFc(-)-G₄-GDF15(N3D):DhCpmFc(+) heterodimers in which the two first polypeptide chains of each heterodimer are linked via an interchain disulfide bond between their respective GDF15 regions.

More particularly, in a specific embodiment, the tetramer comprises:

- (a) two DhCpmFc(+) domains (one each heterodimer) having the sequence of SEQ ID NO:283,
- (b) two DhCpmFc(+) domains (one each heterodimer) having the sequence of SEQ ID NO:48,
- (c) two a GDF15(N3D) polypeptides (one each heterodimer) having the sequence of SEQ ID NO:52, and
- (d) two polypeptide linkers (one each heterodimer) comprising the sequence:

GGGG (SEQ ID NO:58)

each linking the N-terminus of a GDF15(N3D) polypeptide to the C-terminus of a DhCpmFc(-) domain via peptide bonds.

In a preferred embodiment, the first polypeptide comprises the amino acid sequence (linker sequence double underlined):

APELLGGPSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVD
 GVEVHNAKTKPREEQYNSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPA
 PIEKTISKAKGQPREPQVYTLPPSREEMTKNQVSLTCLVKGFYPSDIAVE
 WESNGQPENNYDTPPVLDSDGSFFLYSDLTVDKSRWQQGNVFSQSVME
 ALHNHYTQKSLSLSPGGGGGARDGDHCPLGPGRCRLHTVRASLEDLGWA
 DWVLSPREVQVTMCIGACPSQFRAANMHAQIKTSLHRLKPDTVPAPCCVP
 ASYNPMVLIQKTDGTGVSQTYDDLLAKDCHCI (SEQ ID NO:60),

which is encoded by the nucleic acid sequence:

gcacctgaactcctgggggggaccgtcagtccttcctcttccccccaaaacc
 caaggacaccctcatgatctcccgaccctgaggtcacatgcgtggtgg

tggacgtgagccacgaagaccctgaggtcaagttcaactggtacgtggac
 ggcgtggaggtgcataatgccaaagacaaagccgcgggaggagcagtacaa
 cagcacgtaccgtgtggtcagcgtcctcaccgtcctgcaccaggactggc
 tgaatggcaaggagtacaagtgaagggtctccaacaaagccctcccagcc
 5 cccatcgagaaaaaccatctccaaagccaaagggcagccccgagaaccaca
 ggtgtacaccctgcccccatcccgaggagatgaccaagaaccagggtca
 gcctgacctgcctggtcaaaggcttctatcccagcgacatcgccgtggag
 tgggagagcaatgggcagccggagaacaactacgacaccacgcctcccg
 gctggactccgacggctccttcttctctatagcgacctcaccgtggaca
 10 agagcaggtggcagcaggggaacgtcttctcatgctccgtgatgcatgag
 gctctgcacaaccactacacgcagaagagcctctccctgtctccgggtgg
 aggtggtggagcgcgcgacggagaccactgtccgctcgggccccggcggt
 gctgccgtctgcacacgggtccgcgcgtcgtggaagacctgggctgggccc
 gattgggtgctgtcgccacgggaggtgcaagtgacctgtgcatcggcgc
 15 gtgcccagaccagttccgggcccgaacatgcacgcgcagatcaagacga
 gcctgcaccgcctgaagcccgacacgggtgccagcgccctgctgcgtgccc
 gccagctacaatcccatggtgctcattcaaaagaccgacaccggggtgtc
 gctccagacctatgatgacttggttagccaaagactgccactgcata (SEQ ID NO:59).

In a preferred embodiment, the second polypeptide chain comprises the amino acid
 20 sequence of SEQ ID NO:47, which is encoded by the nucleic acid sequence of SEQ ID
 NO:51.

As discussed above, a tetramer is provided comprising two polypeptide chains having
 the sequence of SEQ ID NO:60 and two polypeptide chains having the sequence of SEQ ID
 NO:47.

25

II.E.6 DhCpmFc(-)-G₄S-GDF15:DhCpmFc(+)

The designation "DhCpmFc(-)-G₄S-GDF15:DhCpmFc(+)" in the instant disclosure
 refers to a heterodimer comprising (i) a first polypeptide chain comprising a GDF15
 polypeptide linked to a DhCpmFc(-) domain via a polypeptide linker comprising the sequence
 30 of SEQ ID NO:61 that connects the N-terminus of the GDF15 polypeptide to the C-terminus
 of the DhCpmFc(-) domain and (ii) a second polypeptide chain comprising a DhCpmFc(+)
 domain.

In certain embodiments, a tetramer is provided, comprising a dimer of two
 DhCpmFc(-)-G₄S-GDF15:DhCpmFc(+) heterodimers in which the two first polypeptide
 35 chains of each heterodimer are linked via an interchain disulfide bond between their
 respective GDF15 regions.

More particularly, in a specific embodiment, the tetramer comprises:

(a) two DhCpmFc(+) domains (one each heterodimer) comprising the sequence of SEQ ID NO:283,

(b) two DhCpmFc(-) domains (one each heterodimer) comprising the sequence of
5 SEQ ID NO:48,

(c) two GDF15 polypeptides (one each heterodimer) comprising the sequence of SEQ ID NO:12, and

(d) two polypeptide linkers (one each heterodimer) comprising the sequence:

GGGGS (SEQ ID NO:61)

10 each linking the N-terminus of a GDF15 polypeptide to the C-terminus of a DhCpmFc(-) domain via peptide bonds.

In a preferred embodiment, the first polypeptide comprises the amino acid sequence (linker double underlined):

APELLGGPSVFLFPPKPKDTLMI SRTPEVTCVVVDVSHEDPEVKFNWYVD
15 GVEVHNAKTKPREEQYNSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPA
PIEKTISKAKGQPREPQVYTLPPSREEMTKNQVSLTCLVKGFYPSDIAVE
WESNGQPENNYDTTPPVLDSDGSFFLYSDLTVDKSRWQQGNVVFSCSVME
ALHNHYTQKSLSLSPGGGGGSARNGDHCPLGPRCCRLHTVRASLEDLGW
ADWVLSPREVQVTMCIGACPSQFRAANMHAQIKTSLHRLKPDTVPAPCCV
20 PASYNPMVLIQKTDGTGVSQTYYDDLLAKDCHCI (SEQ ID NO:63),

which is encoded by the nucleic acid sequence:

gcacctgaactcctggggggaccgtcagtccttcctcttccccccaaaacc
caaggacaccctcatgatctcccggaaccctgaggtcacatgcgtggtgg
tggacgtgagccacgaagaccctgaggtcaagttcaactggtacgtggac
25 ggcgtggaggtgcataatgccaaagacaaagccgcgggaggagcagtacaa
cagcacgtaccgtgtggtcagcgtcctcaccgtcctgcaccaggactggc
tgaatggcaaggagtacaagtgcaaggtctccaacaaagccctcccagcc
cccatcgagaaaaccatctccaaagccaaagggcagccccgagaaccaca
ggtgtacaccctgcccccatcccgaggagatgaccaagaaccagggtca
30 gcctgacctgcctggtcaaaggcttctatcccagcgacatcgccgtggag
tgggagagcaatgggcagccggagaacaactacgacaccacgcctcccgt
gctggactccgacggctccttcttcctctatagcgacctcaccgtggaca
agagcaggtggcagcaggggaacgtcttctcatgctccgtgatgcatgag
gctctgcacaaccactacacgcagaagagcctctccctgtctccgggtgg

aggtggtggatccgcgcgaacggagaccactgtccgctcgggccccgggc
 gttgctgccgtctgcacacgggtccgcgcgtcgtggaagacctgggctgg
 gccgattgggtgctgtcgccacgggaggtgcaagtgaccatgtgcatcgg
 cgcgtgccccgagccagttccgggcggcacaacatgcacgcgcagatcaaga
 5 cgagcctgcaccgcctgaagccccgacacgggtgccagcgccttgcgtgcgtg
 cccgccagctacaatcccatgggtgctcattcaaaagaccgacaccggggt
 gtcgctccagacctatgatgacttgtagccaaagactgccactgcatat
 ga (SEQ ID NO:62) .

In a preferred embodiment, the second polypeptide comprises the amino acid
 10 sequence of SEQ ID NO:47, which is encoded by the nucleic acid sequence of SEQ ID
 NO:51.

As discussed above, a tetramer is provided comprising two polypeptide chains having
 the sequence of SEQ ID NO:63 and two polypeptide chains having the sequence of SEQ ID
 NO:47.

15 *II.E.7 DhCpmFc(-)-(G₄S)₂-GDF15:DhCpmFc(+)*

The designation “DhCpmFc(-)-(G₄S)₂-GDF15:DhCpmFc(+)” in the instant disclosure
 refers to a heterodimer comprising (i) a first polypeptide chain comprising a GDF15
 polypeptide linked to a DhCpmFc(-) domain via a polypeptide linker comprising the sequence
 20 of SEQ ID NO:64 that connects N-terminus of the GDF15 polypeptide to the C-terminus of
 the DhCpmFc(-) domain and (ii) a second polypeptide chain comprising a DhCpmFc(+)
 domain.

In certain embodiments, a tetramer is provided, comprising a dimer of two
 DhCpmFc(-)-(G₄S)₂-GDF15:DhCpmFc(+) heterodimers in which the two first polypeptide
 25 chains of each heterodimer are linked via an interchain disulfide bond between their
 respective GDF15 regions.

More particularly, in a specific embodiment, the tetramer comprises:

- (a) two DhCpmFc(+) domains (one each heterodimer) comprising the sequence of
 SEQ ID NO:283,
- 30 (b) two DhCpmFc(-) domains (one each heterodimer) comprising the sequence of
 SEQ ID NO:48,
- (c) two GDF15 polypeptides (one each heterodimer) comprising the sequence of SEQ
 ID NO:12, and
- (d) two polypeptide linkers (one each heterodimer) comprising the sequence:

35 GGGGSGGGGS (SEQ ID NO:64) ,

each linking the N-terminus of a GDF15 polypeptide to the C-terminus of a DhCpmFc(-) sequence via a peptide bond.

In a preferred embodiment, the first polypeptide chain comprises the amino acid sequence (linker double underlined):

5 APELLGGPSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVD
 GVEVHNAKTKPREEQYNSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPA
 PIEKTIKAKGQPREPQVYTLPPSREEMTKNQVSLTCLVKGFYPSDIAVE
 WESNGQPENNYDTPPVLDSDGSFFLYSDLTVDKSRWQQGNVFSCSVMHE
 ALHNHYTQKSLSLSPGGGGGSGGGGSARNGDHCPLGPGRCCRLHTVRASL
 10 EDLGWADWVLSPREVQVTMCIGACPSQFRAANMHAQIKTSLHRLKPDTVP
 APCCVPASYNPMVLIQKTDGTGVSLSQTYDDLAKDCHCI (SEQ ID NO:66),

which is encoded by the nucleic acid sequence:

gcacctgaactcctggggggaccgtcagtccttcctcttccccccaaaacc
 caaggacaccctcatgatctcccgaccctgaggtcacatgcgtggtgg
 15 tggacgtgagccacgaagaccctgaggtcaagttcaactggtacgtggac
 ggcgtggaggtgcataatgccaaagacaaagccgcgggaggagcagtacaa
 cagcacgtaccgtgtggtcagcgtcctcaccgtcctgcaccaggaactggc
 tgaatggcaaggagtacaagtgcaaggtctccaacaaagccctcccagcc
 cccatcgagaaaaccatctccaaagccaaagggcagccccgagaaccaca
 20 ggtgtacaccctgcccccatcccgaggagatgaccaagaaccaggtca
 gcctgacctgcctggtcaaaggcttctatcccagcgacatcgccgtggag
 tgggagagcaatgggcagccggagaacaactacgacaccacgcctcccgt
 gctggactccgacggctccttcttcctctatagcgacctcaccgtggaca
 agagcaggtggcagcaggggaacgtcttctcatgctccgtgatgcatgag
 25 gctctgcacaaccactacacgcagaagagcctctccctgtctccgggtgg
 aggtggtggatccggaggcgggtggaagcgcgcgcaacggagaccactgtc
 cgctcgggcccggggttgctgccgtctgcacacggtccgcgcgtcgctg
 gaagacctgggctgggcccattgggtgctgtcgccacgggaggtgcaagt
 gaccatgtgcatcggcgcgtgcccgagccagttccgggagggaacatgc
 30 acgcgcagatcaagacgagcctgcaccgcctgaagcccgacacggtgcca
 gcgccctgctgctgcccgccagctacaatcccatggtgctcattcaaaa
 gaccgacaccggggtgtcgctccagacctatgatgacttgtagccaaag
 actgccactgcata (SEQ ID NO:65).

In a preferred embodiment, the second polypeptide chain comprises the amino acid sequence of SEQ ID NO:47, which is encoded by the nucleic acid sequence of SEQ ID NO:51.

As discussed above, a tetramer is provided comprising two polypeptide chains comprising the sequence of SEQ ID NO:66 and two polypeptide chains comprising the sequence of SEQ ID NO:47.

II.E.8 DhCpmFc(-)-(G₄S)₂-GDF15(N3D):DhCpmFc(+)

The designation “DhCpmFc(-)-(G₄S)₂-GDF15(N3D):DhCpmFc(+)” in the instant disclosure refers to a heterodimer comprising (i) a first polypeptide chain comprising a GDF15(N3D) polypeptide linked to a DhCpmFc(-) domain via a linker comprising the sequence of SEQ ID NO:64 that connects the N-terminus of the GDF15(N3D) polypeptide to the C-terminus of the DhCpmFc(-) domain and (ii) a second polypeptide chain comprising a DhCpmFc(+) domain.

In certain embodiments, a tetramer is provided, comprising a dimer of two DhCpmFc(-)-(G₄S)₂-GDF15(N3D):DhCpmFc(+) heterodimers in which the two first polypeptide chains of each heterodimer are linked via an interchain disulfide bond between their respective GDF15 regions.

More particularly, in a specific embodiment, the tetramer comprises:

(a) two DhCpmFc(+) domains (one each heterodimer) comprising the sequence of SEQ ID NO:283,

(b) two DhCpmFc(-) domains (one each heterodimer) comprising the sequence of SEQ ID NO:48,

(c) two GDF15(N3D) polypeptides (one each heterodimer) comprising the sequence of SEQ ID NO:52, and

(d) two polypeptide linkers (one each heterodimer) comprising the sequence of SEQ ID NO:64, each linking the N-terminus of a GDF15(N3D) polypeptide to the C-terminus of a DhCpmFc(-) domain via a peptide bond.

In a preferred embodiment, the first polypeptide chain comprises the amino acid sequence (linker sequence double underlined):

APELLGGPSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVD
GVEVHNAKTKPREEQYNSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPA
PIEKTISKAKGQPREPQVYTLPPSREEMTKNQVSLTCLVKGFYPSDIAVE
WESNGQPENNYDTTPPVLDSDGSFFLYSDLTVDKSRWQQGNVFSQSVME
ALHNHYTQKSLSLSPGGGGGSGGGGSARDGDHCPLGPGRCCRLHTVRASL

EDLGWADWVLSPREVQVTCIGACPSQFRAANMHAQIKTSLHRLKPDTVP
APCCVPASYNPMVLIQKTDGTGVSLLQTYDDLAKDCHCI (SEQ ID NO:68),

which is encoded by the nucleic acid sequence:

gcacctgaactcctggggggaccgtcagtccttctcttccccccaaaacc
5 caaggacaccctcatgatctcccgaccctgaggtcacatgcgtggtgg
tggacgtgagccacgaagaccctgaggtcaagttcaactggtacgtggac
ggcgtggaggtgcataatgccaaagacaaagccgcgggaggagcagtacaa
cagcacgtaccgtgtggtcagcgtcctcaccgtcctgcaccaggactggc
tgaatggcaaggagtacaagtgaaggtctccaacaaagccctcccagcc
10 cccatcgagaaaaccatctccaaagccaaagggcagccccgagaaccaca
ggtgtacaccctgcccccatcccgaggagatgaccaagaaccagggtca
gcctgacctgcctggtcaaaggcttctatcccagcgacatcgccgtggag
tgggagagcaatgggcagccggagaacaactacgacaccacgcctcccg
gctggactccgacggctccttcttctctatagcgacctcaccgtggaca
15 agagcaggtggcagcaggggaacgtcttctcatgctccgtgatgcatgag
gctctgcacaaccactacacgcagaagagcctctccctgtctccgggtgg
aggtggtggatccggaggcgggtggaagcgcgcgcgacggagaccactgtc
cgctcggggcccgggcgttgctgccgtctgcacacgggtccgcgcgtcgctg
gaagacctgggctgggcccattgggtgctgtcgccacgggaggtgcaagt
20 gaccatgtgcatcggcgcgtgcccagagccagttccggggcggaacatgc
acgcgcagatcaagacgagcctgcaccgcctgaagcccgcacgggtgcca
gcgccctgctgctgcccgcagctacaatcccattggtgctcattcaaaa
gaccgacaccggggtgtcgctccagacctatgatgacttgtagccaaag
actgccactgcata (SEQ ID NO:67).

25 In a preferred embodiment, the second polypeptide chain comprises the amino acid sequence of SEQ ID NO:47, which is encoded by the nucleic acid sequence of SEQ ID NO:51.

As discussed above, a tetramer is provided comprising two polypeptide chains having the sequence of SEQ ID NO:68 and two polypeptide chains having the sequence of SEQ ID
30 NO:47.

II.E.9 DhCpmFc(-)-G₄P-GDF15:DhCpmFc(+)

The designation “DhCpmFc(-)-(G₄P)-GDF15:DhCpmFc(+)” in the instant disclosure refers to a heterodimer comprising (i) a first polypeptide chain comprising a GDF15

polypeptide linked to a DhCpmFc(-) domain via a linker comprising the sequence of SEQ ID NO:69 that connects the N-terminus

of the GDF15 polypeptide to the C-terminus of the DhCpmFc(-) domain and (ii) a second polypeptide chain comprising a DhCpmFc(+) domain.

- 5 In certain embodiments, a tetramer is provided, comprising a dimer of two DhCpmFc(-)-(G₄P)-GDF15:DhCpmFc(+) heterodimers in which the two first polypeptide chains of each heterodimer are linked via an interchain disulfide bond between their respective GDF15 regions.

More particularly, in a specific embodiment, the tetramer comprises:

- 10 (a) two DhCpmFc(+) domains (one each heterodimer) comprising the sequence of SEQ ID NO:283,

(b) two DhCpmFc(-) domains (one each heterodimer) comprising the sequence of SEQ ID NO:48,

- 15 (c) two GDF15 polypeptides (one each heterodimer) comprising the sequence of SEQ ID NO:12, and

(d) two polypeptide linkers (one each heterodimer) comprising the sequence:

GGGGP (SEQ ID NO:69),

each linking the N-terminus of a GDF15 polypeptide to the C-terminus of a DhCpmFc(-) domain via a peptide bond.

- 20 In a preferred embodiment, the first polypeptide chain comprises the amino acid sequence (linker sequence double underlined):

APELLGGPSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVD
GVEVHNAKTKPREEQYNSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPA
PIEKTISKAKGQPREPQVYTLPPSREEMTKNQVSLTCLVKGFYPSDIAVE
25 WESNGQPENNYDTPPVLDSDGSFFLYSDLTVDKSRWQQGNVFSQSVME
ALHNHYTQKSLSLSPGGGGGPARNGDHCPLGPGRCRLHTVRASLEDLGW
ADWVLSPREVQVTMCIGACPSQFRAANMHAQIKTSLHRLKPDTVPAPCCV
PASYNPMVLIQKTDGTGVSQTYYDILLAKDCHCI (SEQ ID NO:71),

which is encoded by the nucleic acid sequence:

- 30 gcacctgaactcctggggggaccgtcagtccttcctcttccccccaaaacc
caaggacaccctcatgatctcccgaccctgaggtcacatgcgtggtgg
tggacgtgagccacgaagaccctgaggtcaagttcaactggtacgtggac
ggcgtggaggtgcataatgccaaagacaaagccgaggaggagcagtacaa
cagcacgtaccgtgtggtcagcgtcctcaccgtcctgcaccaggactggc

tgaatggcaaggagtacaagtgaaggtctccaacaaagccctcccagcc
 cccatcgagaaaaaccatctccaaagccaaagggcagccccgagaaccaca
 ggtgtacaccctgcccccatccccgggaggagatgaccaagaaccaggtca
 gcctgacctgcctggtcaaaggettctatcccagcgacatcgccgtggag
 5 tgggagagcaatgggcagccggagaacaactacgacaccacgcctcccgt
 gctggactccgacggctccttcttctctatagcgacctcacctgggaca
 agagcaggtggcagcaggggaacgtcttctcatgctccgtgatgcatgag
 gctctgcacaaccactacacgcagaagagcctctccctgtctccgggtgg
 aggtggtggacccgcgcgaacggagaccactgtccgctcgggccccgggc
 10 gttgctgccgtctgcacacggtccgcgcgtcgctggaagacctgggctgg
 gccgattgggtgctgtcgccacgggaggtgcaagtgaccatgtgcatcgg
 cgcgtgccccgagccagttccgggcggaacatgcacgcgcagatcaaga
 cgagcctgcaccgcctgaagccccgacacggtgccagcgccctgctgcgtg
 ccgcccagctacaatcccatggtgctcattcaaaagaccgacaccgggt
 15 gtcgctccagacctatgatgacttgtagccaaagactgccactgcata (SEQ ID
 NO:70) .

In a preferred embodiment, the second polypeptide chain comprises the amino acid sequence of SEQ ID NO:47, which is encoded by the nucleic acid sequence of SEQ ID NO:51.

20 As discussed above, a tetramer is provided comprising two polypeptide chains having the sequence of SEQ ID NO:71 and two polypeptide chains having the sequence of SEQ ID NO:47.

II.E.10 DhCpmFc(-)-(G₄P)₂-GDF15:DhCpmFc(+)

25 The designation “DhCpmFc(-)-(G₄P)₂-GDF15:DhCpmFc(+)” in the instant disclosure refers to a heterodimer comprising (i) a first polypeptide chain comprising a GDF15 polypeptide linked to a DhCpmFc(-) domain via a linker comprising the sequence of SEQ ID NO:72 that connects the N-terminus of the GDF15 polypeptide to the C-terminus of the DhCpmFc(-) domain and (ii) a second v comprising a DhCpmFc(+) sequence.

30 In certain embodiments, a tetramer is provided, comprising a dimer of two DhCpmFc(-)-(G₄P)₂-GDF15:DhCpmFc(+) heterodimers in which the two first polypeptide chains of each heterodimer are linked via an interchain disulfide bond between their respective GDF15 regions.

More particularly, in a specific embodiment, the tetramer comprises:

35 (a) two DhCpmFc(+) domains (one each heterodimer) comprising the sequence of SEQ ID NO:283,

(b) two DhCpmFc(-) domains (one each heterodimer) comprising the sequence of SEQ ID NO:48,

(c) two GDF15 polypeptides (one each heterodimer) comprising the sequence of SEQ ID NO:12, and

5 (d) two polypeptide linkers (one each heterodimer) comprising the sequence:

GGGGPGGGGP (SEQ ID NO:72),

each linking the N-terminus of a GDF15 polypeptide to the C-terminus of a DhCpmFc(-) domain via a peptide bond.

10 In a preferred embodiment, the first polypeptide chain comprises the amino acid sequence (linker sequence double underlined):

APELLGGPSVFLFPPKPKDTLMI SRTPEVTCVVVDVSHEDPEVKFNWYVD
GVEVHNAKTKPREEQYNSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPA
PIEKTISKAKGQPREPQVYTLPPSREEMTKNQVSLTCLVKGFYPSDIAVE
WESNGQPENNYDTPPVLDSDGSFFLYSDLTVDKSRWQQGNVFSQSVME
15 ALHNHYTQKSLSLSPGGGGGPGGGGPARNGDHCPLGPGRCCRLHTVRASL
EDLGWADWVLSPREVQVTMCIGACPSQFRAANMHAQIKTSLHRLKPDTVP
APCCVPASYNPMVLIQKTDGTGVS LQTYDDLAKDCHCI (SEQ ID NO:74),

which is encoded by the nucleic acid sequence:

gcacctgaactcctggggggaccgtcagtccttcctcttccccccaaaacc
20 caaggacaccctcatgatctcccgaccctgaggtcacatgcgtggtgg
tggacgtgagccacgaagaccctgaggtcaagttcaactggtacgtggac
ggcgtggaggtgcataatgccaaagacaaagccgcggaggagcagtacaa
cagcacgtaccgtgtggtcagcgtcctcaccgtcctgcaccaggactggc
tgaatggcaaggagtacaagtgaaggtctccaacaaagccctcccagcc
25 cccatcgagaaaaccatctccaaagccaaagggcagccccgagaaccaca
ggtgtacaccctgcccccatcccgaggagatgaccaagaaccaggtca
gcctgacctgcctggtcaaaggcttctatcccagcgacatcgccgtggag
tgggagagcaatgggcagccggagaacaactacgacaccacgcctcccgt
gctggactccgacggctccttcttctctatagcgacctcaccgtggaca
30 agagcaggtggcagcaggggaacgtcttctcatgctccgtgatgcatgag
gctctgcacaaccactacacgcagaagagcctctccctgtctccgggtgg
aggtggtggacctggaggcgtggaccagcgcgcaacggagaccactgtc
cgctcggggcccggttgctgccgtctgcacacgggtccgcgcgtcgctg
gaagacctgggctgggcccattgggtgctgtcgccacgggaggtgcaagt

gaccatgtgcatcgggcgctgccccgagccagttccgggcggaacatgc
 acgcgcagatcaagacgagcctgcaccgcctgaagcccgacacggtgcca
 ggcgcctgctgctgccccgccagctacaatcccatgggtgctcattcaaaa
 gaccgacaccgggggtgctcgctccagacctatgatgacttgtagccaaag
 5 actgccactgcatatga (SEQ ID NO:73).

In a preferred embodiment, the second polypeptide chain comprises the amino acid sequence of SEQ ID NO:47, which is encoded by the nucleic acid sequence of SEQ ID NO:51.

As discussed above, a tetramer is provided comprising two polypeptide chains having
 10 the sequence of SEQ ID NO:74 and two polypeptide chains comprising the sequence of SEQ ID NO:47.

II.E.11 DhCpmFc(-)-G₄Q-GDF15:DhCpmFc(+)

The designation “DhCpmFc(-)-G₄Q-GDF15:DhCpmFc(+)” in the instant disclosure
 15 refers to a heterodimer comprising (i) a first polypeptide chain comprising a GDF15 polypeptide linked to a DhCpmFc(-) domain via a linker comprising the sequence of SEQ ID NO:75 that connects the N-terminus of the GDF15 polypeptide to the C-terminus of the DhCpmFc(-) domain and (ii) a second polypeptide chain comprising a DhCpmFc(+) domain.

In certain embodiments, a tetramer is provided, comprising a dimer of two
 20 DhCpmFc(-)-G₄Q-GDF15:DhCpmFc(+) heterodimers in which the two first polypeptide chains of each heterodimer are linked via an interchain disulfide bond between their respective GDF15 regions.

More particularly, in a specific embodiment, the tetramer comprises:

(a) two DhCpmFc(+) chains (one each heterodimer) comprising the sequence of SEQ
 25 ID NO:283,

(b) two DhCpmFc(-) chains (one each heterodimer) comprising the sequence of SEQ ID NO:48,

(c) two GDF15 polypeptides (one each heterodimer) comprising the sequence of SEQ ID NO:12, and

30 (d) two polypeptide linkers (one each heterodimer) comprising the sequence:

GGGGQ (SEQ ID NO:75)

each linking the N-terminus of a GDF15 polypeptide to the C-terminus of a DhCpmFc(-) domain via a peptide bond.

In a preferred embodiment, the first polypeptide chain comprises the amino acid
 35 sequence (linker sequence double underlined):

APELLGGPSVFLFPPKPKDTLMISRTPEVTCVVDVSHEDPEVKFNWYVD
 GVEVHNAKTKPREEQYNSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPA
 PIEKTISKAKGQPREPQVYTLPPSREEMTKNQVSLTCLVKGFYPSDIAVE
 WESNGQPENNYDTPPVLDSDGSFFLYSDLTVDKSRWQQGNVFSQVMHE
 5 ALHNHYTQKSLSLSPGGGGQARNGDHCPLGPGRCCRLHTVRASLEDLGW
 ADWVLSPREVQVTMCIGACPSQFRAANMHAQIKTSLHRLKPDTPAPCCV
 PASYNPMVLIQKTDGTGSLQTYDDLLAKDCHCI (SEQ ID NO:77),

which is encoded by the nucleic acid sequence:

gcacctgaactcctggggggaccgtcagtccttcctcttccccccaaaacc
 10 caaggacaccctcatgatctcccgaccctgaggtcacatgcgtggtgg
 tggacgtgagccacgaagaccctgaggtcaagttcaactggtacgtggac
 ggcgtggaggtgcataatgccaaagacaaagccgcgggaggagcagtacaa
 cagcacgtaccgtgtggtcagcgtcctcaccgtcctgcaccaggactggc
 tgaatggcaaggagtacaagtgaaggtctccaacaaagccctcccagcc
 15 cccatcgagaaaaccatctccaaagccaaagggcagccccgagaaccaca
 ggtgtacaccctgcccccatcccgaggagatgaccaagaaccagggtca
 gcctgacctgcctggtcaaaggcttctatcccagcgacatcgccgtggag
 tgggagagcaatgggcagccggagaacaactacgacaccacgcctcccg
 gctggactccgacggctccttcttctctatagcgacctcactgtggaca
 20 agagcaggtggcagcaggggaacgtcttctcatgctccgtgatgcag
 gctctgcacaaccactacacgcagaagagcctctccctgtctccgggtgg
 aggtggtggacaggcgcgcaacggagaccactgtccgctcgggccccggc
 gttgctgccgtctgcacacggtccgcgcgtcgtggaagacctgggctgg
 gccgattgggtgctgtcgccacgggaggtgcaagtgacctgtgcatcgg
 25 cgcgtgcccagaccagttccgggcggaacatgcacgcgcagatcaaga
 cgagcctgcaccgcctgaagcccgcacacgggtgccagcgccctgctgcgtg
 cccgccagctacaatcccatggtgctcattcaaaagaccgacaccggggt
 gtcgctccagacctatgatgacttgtagccaaagactgccactgcatat
 ga (SEQ ID NO:76).

30 In a preferred embodiment, the second polypeptide chain comprises the amino acid
 sequence of SEQ ID NO:47, which is encoded by the nucleic acid sequence of SEQ ID
 NO:51.

As discussed above, a tetramer is provided comprising two polypeptide chains having
 the sequence of SEQ ID NO:77 and two polypeptide chains having the sequence of SEQ ID
 35 NO:47.

II.E.12 DhCpmFc(-)-(G₄Q)₂-GDF15:DhCpmFc(+)

The designation “DhCpmFc(-)-(G₄Q)₂-GDF15:DhCpmFc(+)” in the instant disclosure refers to a heterodimer comprising (i) a first polypeptide chain comprising a GDF15 polypeptide linked to a DhCpmFc(-) domain via a linker comprising the sequence of SEQ ID NO:78 that connects the N-terminus of the GDF15 polypeptide to the C-terminus of the DhCpmFc(-) domain and (ii) a second polypeptide chain comprising a DhCpmFc(+) domain.

In certain embodiments, a tetramer is provided, comprising a dimer of two DhCpmFc(-)-(G₄Q)₂-GDF15:DhCpmFc(+) heterodimers in which the two first polypeptide chains of each heterodimer are linked via an interchain disulfide bond between their respective GDF15 regions.

More particularly, in a specific embodiment, the tetramer comprises:

(a) two DhCpmFc(+) domains (one each heterodimer) comprising the sequence of SEQ ID NO:283,

(b) two DhCpmFc(-) domains (one each heterodimer) comprising the sequence of SEQ ID NO:48,

(c) two GDF15 polypeptides (one each heterodimer) comprising the sequence of SEQ ID NO:12, and

(d) two polypeptide linkers (one each heterodimer) comprising the sequence:

GGGGQGGGGQ (SEQ ID NO:78)

each linking the N-terminus of a GDF15 polypeptide to the C-terminus of a DhCpmFc(-) domain via a peptide bond.

In a preferred embodiment, the first polypeptide chain comprises the amino acid sequence (linker sequence double underlined):

APELLGGPSVFLFPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVD
 GVEVHNAKTKPREEQYNSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPA
 PIEKTISKAKGQPREPQVYTLPPSREEMTKNQVSLTCLVKGFYPSDIAVE
 WESNGQPENNYDTTPPVLDSDGSFFLYSDLTVDKSRWQQGNVFSQSVME
 30 ALHNHYTQKSLSLSPGGGGQGGGGQARNGDHCPLGPRCCRLHTVRASL
 EDLGWADWVLSPREVQVTMCIGACPSQFRAANMHAQIKTSLHRLKPDTPV
 APCCVPASYNPMVLIQKTDGTGVSLLQTYDDLLAKDCHCI (SEQ ID NO:80),

which is encoded by the nucleic acid sequence:

gcacctgaactcctggggggaccgtcagtccttctcttccccccaaaacc
 caaggacaccctcatgatctcccgaccctgaggtcacatgcgtggtgg
 tggacgtgagccacgaagaccctgaggtcaagttcaactggtacgtggac
 ggcgaggaggtgcataatgccaaagacaaagccgcgggaggagcagtacaa
 5 cagcacgtaccgtgtggtcagcgtcctcaccgtcctgcaccaggactggc
 tgaatggcaaggagtacaagtgcaaggtctccaacaaagccctcccagcc
 cccatcgagaaaaccatctccaaagccaaagggcagccccgagaaccaca
 ggtgtacaccctgcccccatcccgaggagatgaccaagaaccaggtca
 gcctgacctgcctggtcaaaggcttctatcccagcgacatcgccgtggag
 10 tgggagagcaatgggcagccggagaacaactacgacaccacgcctcccgt
 gctggactccgacggctccttcttctctatagcgacctcaccgtggaca
 agagcaggtggcagcaggggaacgtcttctcatgctccgtgatgcatgag
 gctctgcacaaccactacacgcagaagagcctctccctgtctccgggtgg
 aggtggtggacagggaggcgggtggacagggcgcgaacggagaccactgtc
 15 cgctcggggcccgggcggttctgctgccgtctgcacacgggtccgcgcgtcgtg
 gaagacctgggctgggcccattgggtgctgtcgccacgggaggtgcaagt
 gaccatgtgcatcggcgcgtgcccagagccagttccggggcggaacatgc
 acgcgcagatcaagacgagcctgcaccgcctgaagcccgcacaggtgcca
 ggcacctgctgctgcccgcagctacaatcccattggtgctcattcaaaa
 20 gaccgacaccggggtgtcgctccagacctatgatgacttgtagccaaag
 actgccactgcatatga (SEQ ID NO:79).

In a preferred embodiment, the second polypeptide chain comprises the amino acid sequence of SEQ ID NO:47, which is encoded by the nucleic acid sequence of SEQ ID NO:51.

25 As discussed above, a tetramer is provided comprising two polypeptide chains having the sequence of SEQ ID NO:80 and two polypeptide chains having the sequence of SEQ ID NO:47.

II.E.13 DhCpmFc(-)-(G₄Q)₂-GDF15(N3D):DhCpmFc(+)

30 The designation “DhCpmFc(-)-(G₄Q)₂-GDF15(N3D):DhCpmFc(+)” in the instant disclosure refers to a heterodimer comprising (i) a first polypeptide chain comprising a GDF15(N3D) polypeptide linked to a DhCpmFc(-) domain via a linker comprising the sequence of SEQ ID NO:78 that connects the N-terminus of the GDF15(N3D) polypeptide to the C-terminus of the DhCpmFc(-) domain and (ii) a second polypeptide chain comprising a
 35 DhCpmFc(+) domain.

In certain embodiments, a tetramer is provided, comprising a dimer of two DhCpmFc(-)-(G₄Q)₂-GDF15(N3D):DhCpmFc(+) heterodimers in which the two first polypeptide chains of each heterodimer are linked via an interchain disulfide bond between their respective GDF15 regions.

5 More particularly, in a specific embodiment, the tetramer comprises:

(a) two DhCpmFc(+) domains (one each heterodimer) comprising the sequence of SEQ ID NO:283,

(b) two DhCpmFc(-) domains (one each heterodimer) comprising the sequence of SEQ ID NO:48,

10 (c) two GDF15(N3D) polypeptides (one each heterodimer) comprising the sequence of SEQ ID NO:52, and

(d) two polypeptide linkers (one each heterodimer) comprising the sequence of SEQ ID NO:78 each linking the N-terminus of a GDF15(N3D) polypeptide to the C-terminus of a DhCpmFc(-) domain via a peptide bond.

15 In a preferred embodiment, the first polypeptide chain comprises the amino acid sequence (linker sequence double underlined):

APELLGGPSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVD
GVEVHNAKTKPREEQYNSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPA
PIEKTISKAKGQPREPQVYTLPPSREEMTKNQVSLTCLVKGFYPSDIAVE
20 WESNGQPENNYDTPPVLDSDGSFFLYSDLTVDKSRWQQGNVFSCSVMHE
ALHNHYTQKSLSLSPGGGGQGGGGQARDGDHCPLGPRCCRLHTVRASL
EDLGWADWVLSPREVQVTMCIGACPSQFRAANMHAQIKTSLHRLKPDTPV
APCCVPASYNPMVLIQKTDGTGVSLLQTYDDLAKDCHCI (SEQ ID NO:82),

which is encoded by the nucleic acid sequence:

25 gcacctgaactcctggggggaccgtcagtccttcctcttccccccaaaacc
caaggacaccctcatgatctcccgaccctgaggtcacatgcgtggtgg
tggacgtgagccacgaagaccctgaggtcaagttcaactggtacgtggac
ggcgtggaggtgcataatgccaaagacaaagccgcggaggagcagtacaa
cagcacgtaccgtgtggtcagcgtcctcaccgtcctgcaccaggactggc
30 tgaatggcaaggagtacaagtgaaggtctccaacaaagccctcccagcc
cccatcgagaaaaccatctccaaagccaaagggcagccccgagaaccaca
ggtgtacaccctgcccccatcccgaggagatgaccaagaaccagggtca
gcctgacctgcctggtcaaaggcttctatcccagcgacatcgccgtggag
tgggagagcaatgggcagccggagaacaactacgacaccacgcctcccg
35 gctggactccgacggctccttcttctctatagcgacctcaccgtggaca

agagcaggtggcagcaggggaacgtcttctcatgctccgtgatgcatgag
 gctctgcacaaccactacacgcagaagagcctctccctgtctccgggtgg
 aggtggtggacagggaggcgggtggacagggcgcgcgacggagaccactgtc
 cgctcggggcccgggcggttgctgcegtctgcacacgggtccgcgcgctcgctg
 5 gaagacctgggctgggcccattgggtgctgtcgccacgggaggtgcaagt
 gaccatgtgcatcggcgcgtgcccagagccagttccggggcggaacatgc
 acgcgcagatcaagacgagcctgcaccgcctgaagcccgacacggtgcca
 gcgcctgtgtgctgcccgcagctacaatcccattggtgtcattcaaaa
 gaccgacaccggggtgtcgctccagacctatgatgacttgtagccaaag
 10 actgccactgcata (SEQ ID NO:81).

In a preferred embodiment, the second polypeptide chain comprises the amino acid sequence of SEQ ID NO:47, which is encoded by the nucleic acid sequence of SEQ ID NO:51.

As discussed above, a tetramer is provided comprising two polypeptide chains having the sequence of SEQ ID NO:82 and two polypeptide chains having the sequence of SEQ ID NO:47.

II.E.14 DhCpmFc(-)-(G₄Q)₂-GDF15(Ndel3):DhCpmFc(+)

The designation “DhCpmFc(-)-(G₄Q)₂-GDF15(Ndel3):DhCpmFc(+)” in the instant disclosure refers to a heterodimer comprising (i) a first polypeptide chain comprising a GDF15(Ndel3) polypeptide linked to a DhCpmFc(-) domain via a linker comprising the sequence of SEQ ID NO:78 that connects the N-terminus of the GDF15(Ndel3) polypeptide to the C-terminus of the DhCpmFc(-) domain and (ii) a second polypeptide chain comprising a DhCpmFc(+) domain.

In certain embodiments, a tetramer is provided, comprising a dimer of two DhCpmFc(-)-(G₄Q)₂-GDF15(Ndel3):DhCpmFc(+) heterodimers in which the two first polypeptide chains of each heterodimer are linked via an interchain disulfide bond between their respective GDF15 regions.

More particularly, in a specific embodiment, the tetramer comprises:

(a) two DhCpmFc(+) domains (one each heterodimer) comprising the sequence of SEQ ID NO:283,

(b) two DhCpmFc(-) domains (one each heterodimer) comprising the sequence of SEQ ID NO:48,

(c) two GDF15(Ndel3) polypeptides (one each heterodimer) comprising the sequence of SEQ ID NO:55, and

(d) two polypeptide linkers (one each heterodimer) comprising the sequence of SEQ ID NO:78 each linking the N-terminus of a GDF15(Ndel3) polypeptide to the C-terminus of a DhCpmFc(-) domain via a peptide bond.

In a preferred embodiment, the first polypeptide chain comprises the amino acid
5 sequence (linker sequence double underlined):

APELLGGPSVFLFPPKPKDTLMI SRTPEVTCVVVDVSHEDPEVKFKNWYVD
GVEVHNAKTKPREEQYNSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPA
PIEKTISKAKGQPREPQVYTLPPSREEMTKNQVSLTCLVKGFYPSDIAVE
WESNGQPENNYDTPPVLDSDGSFFLYSDLTVDKSRWQQGNVFCFSVMHE
10 ALHNHYTQKSLSLSPGGGGGQGGGGQGDHCP LGPGRCCRLHTVRASLEDL
GWADWVLSPREVQVTMCIGACPSQFRAANMHAQIKTSLHRLKPDTPAPC
CVPASYNPMVLIQKTD TGVS LQTYDDLLAKDCHCI (SEQ ID NO:84),

which is encoded by the nucleic acid sequence:

gcacctgaactcctggggggaccgtcagtccttctctccccccaaaacc
15 caaggacaccctcatgatctcccgaccctgaggtcacatgcgtggtgg
tggacgtgagccacgaagaccctgaggtcaagttcaactggtacgtggac
ggcgtggaggtgcataatgccaagacaaagccgcgggaggagcagtacaa
cagcacgtaccgtgtggtcagcgtcctcaccgtcctgcaccaggactggc
tgaatggcaaggagtacaagtgaaggtctccaacaaagccctcccagcc
20 cccatcgagaaaaccatctccaaagccaaagggcagccccgagaaccaca
ggtgtacaccctgcccccatcccgaggagatgaccaagaaccagggtca
gcctgacctgcctggtcaaaggcttctatcccagcgacatcgccgtggag
tgggagagcaatgggcagccggagaacaactacgacaccacgcctcccgt
gctggactccgacggctccttcttctctatagcgacctcaccgtggaca
25 agagcaggtggcagcaggggaacgtcttctcatgctccgtgatgcatgag
gctctgcacaaccactacacgcagaagagcctctccctgtctccgggtgg
aggtggtggacagggaggcgggtggacagggagaccactgtccgctcgggc
ccgggcgttgctgccgtctgcacacgggtccgcgcgtcgctggaagacctg
ggctgggcccattgggtgctgtcgccacgggaggtgcaagtgaccatgtg
30 catcggcgcgtgcccagccagttccgggcggcaaacatgcacgcgcaga
tcaagacgagcctgcaccgcctgaagcccagacaggtgccagcgcctgc
tgcgtgcccgcagctacaatcccatggtgctcattcaaaagaccgacac
cgggggtgtcgctccagacctatgatgacttgtagccaaagactgccact
gcata (SEQ ID NO:83).

In a preferred embodiment, the second polypeptide chain comprises the amino acid sequence of SEQ ID NO:47, which is encoded by the nucleic acid sequence of SEQ ID NO:51.

As discussed above, a tetramer is provided comprising two polypeptide chains having the sequence of SEQ ID NO:84 and two polypeptide chains having the sequence of SEQ ID NO:47.

II.F. Charged pair (delHinge) Cysteine Clamp

A “cysteine clamp” mutation may be introduced into a Fc domain, such as a CpmFc(+) domain, a CpmFc(-) domain, a DhCpmFc(+) domain, or a DhCpmFc(-) domain. A “cysteine clamp” mutation typically involves the introduction of a cysteine into the CH3 domain of an Fc domain at a specific location through mutation so that when incubated with another Fc domain, also having a cysteine introduced into the CH3 domain at a specific location through mutation, a disulfide bond (cysteine clamp) may be formed between the two Fc domains (e.g., between a CpmFc(+) domain having a “cysteine clamp” mutation and a CpmFc(-) domain having a “cysteine clamp” mutation or between a DhCpmFc(+) domain having a “cysteine clamp” mutation and a DhCpmFc(-) domain having a “cysteine clamp” mutation). An Fc domain may contain one or more such cysteine clamp mutations.

In one embodiment, a cysteine clamp is provided by introducing a serine to cysteine mutation (S354C) into a first Fc domain and a tyrosine to cysteine mutation (Y349C) into a second Fc domain.

The designation “DhCpmFc(-)(S354C)” domain in the instant disclosure refers to an DhCpmFc(-) domain comprising a serine to cysteine mutation (S354C). The designation “DhCpmFc(+)(S354C)” domain in the instant disclosure refers to an DhCpmFc(+) domain comprising a serine to cysteine mutation (S354C). The designation “DhCpmFc(-)(Y349C)” domain in the instant disclosure refers to an DhCpmFc(-) domain comprising a serine to cysteine mutation (Y349C). The designation “DhCpmFc(+)(Y349C)” domain in the instant disclosure refers to an DhCpmFc(+) domain comprising a serine to cysteine mutation (Y349C).

The designation “CpmFc(-)(S354C)” domain in the instant disclosure refers to an CpmFc(-) domain comprising a serine to cysteine mutation (S354C). The designation “CpmFc(+)(S354C)” domain in the instant disclosure refers to an CpmFc(+) domain comprising a serine to cysteine mutation (S354C). The designation “CpmFc(-)(Y349C)” domain in the instant disclosure refers to an CpmFc(-) domain comprising a serine to cysteine mutation (Y349C). The designation “CpmFc(+)(Y349C)” domain in the instant disclosure refers to an CpmFc(+) domain comprising a serine to cysteine mutation (Y349C).

In another embodiment, a cysteine clamp is provided by introducing a leucine to cysteine mutation (L351C) into both a first and Fc domain.

The designation “DhCpmFc(-)(L351C)” domain in the instant disclosure refers to an DhCpmFc(-) domain comprising a serine to cysteine mutation (L351C). The designation
5 “DhCpmFc(+)(L351C)” domain in the instant disclosure refers to an DhCpmFc(+) domain comprising a serine to cysteine mutation (L351C).

The designation “CpmFc(-)(L351C)” domain in the instant disclosure refers to an CpmFc(-) domain comprising a serine to cysteine mutation (L351C). The designation
10 “CpmFc(+)(L351C)” domain in the instant disclosure refers to an CpmFc(+) domain comprising a serine to cysteine mutation (L351C).

The C-terminal lysine (K447) optionally may be deleted in the CpmFc(+) domain, the CpmFc(-) domain, or both. This may be advantageous, for example, when a peptide is fused at the C-terminus to reduce proteolysis of the fusion protein.

In some embodiments a heterodimer is provided comprises (i) a first polypeptide
15 chain comprising a GDF15 region linked to a CpmFc(+) domain comprising a cysteine clamp mutation and (ii) a second polypeptide chain comprising a CpmFc(-) domain comprising a cysteine clamp mutation. In some embodiments, the N-terminus of the GDF15 region is linked to the C-terminus of the CpmFc(+) domain comprising a cysteine clamp mutation, directly or via a polypeptide linker. In other embodiments, the N-terminus of the CpmFc(+)
20 domain comprising a cysteine clamp mutation is linked to the C-terminus of the GDF15 region, directly or via a polypeptide linker.

In some embodiments a heterodimer is provided comprises (i) a first polypeptide chain comprising a GDF15 region linked to a CpmFc(-) domain comprising a cysteine clamp mutation and (ii) a second polypeptide chain comprising a CpmFc(+) domain comprising a
25 cysteine clamp mutation. In some embodiments, the N-terminus of the GDF15 region is linked to the C-terminus of the CpmFc(-) domain comprising a cysteine clamp mutation, directly or via a polypeptide linker. In other embodiments, the N-terminus of the CpmFc(-) domain comprising a cysteine clamp mutation is linked to the C-terminus of the GDF15 region, directly or via a polypeptide linker.

30 In some embodiments a heterodimer is provided comprises (i) a first polypeptide chain comprising a GDF15 region linked to a DhCpmFc(+) domain comprising a cysteine clamp mutation and (ii) a second polypeptide chain comprising a DhCpmFc(-) domain comprising a cysteine clamp mutation. In some embodiments, the N-terminus of the GDF15 region is linked to the C-terminus of the DhCpmFc(+) domain comprising a cysteine clamp
35 mutation, directly or via a polypeptide linker. In other embodiments, the N-terminus of the DhCpmFc(+) domain comprising a cysteine clamp mutation is linked to the C-terminus of the GDF15 region, directly or via a polypeptide linker.

In some embodiments a heterodimer is provided comprises (i) a first polypeptide chain comprising a GDF15 region linked to a DhCpmFc(-) domain comprising a cysteine clamp mutation and (ii) a second polypeptide chain comprising a DhCpmFc(+) domain comprising a cysteine clamp mutation. In some embodiments, the N-terminus of the GDF15 region is linked to the C-terminus of the DhCpmFc(-) domain comprising a cysteine clamp mutation, directly or via a polypeptide linker. In other embodiments, the N-terminus of the DhCpmFc(-) domain comprising a cysteine clamp mutation is linked to the C-terminus of the GDF15 region, directly or via a polypeptide linker.

In some embodiments, a tetramer is provided comprising a dimer of two such heterodimers in which the two first polypeptide chains of the heterodimers are linked via an interchain disulfide bond between their respective GDF15 regions. See Figure 5 for a graphic depiction of an embodiment of a tetramer comprising two heterodimers, where each heterodimer comprises (i) a first polypeptide chain comprising a GDF15 polypeptide linked via a polypeptide linker to a DhCpmFc(+)(L351C) domain and (ii) a second polypeptide chain comprising a DhCpmFc(-)(L351C) domain.

II.F.1 DhCpmFc(+)(S354C)-GDF15(N3D):DhCpmFc(-)(Y349C)

The designation “DhCpmFc(+)(S354C)-GDF15(N3D):DhCpmFc(-)(Y349C)” in the instant disclosure refers to a heterodimer comprising (i) a first polypeptide chain comprising a GDF15(N3D) polypeptide, the N-terminus of which is linked directly to the C-terminus of a DhCpmFc(+)(S354C) domain, and (ii) a second polypeptide chain comprising a DhCpmFc(-)(Y349C) domain. The cysteine clamp mutations allow the first and second polypeptide chains to be linked via an interchain disulfide bond between C354 of the first polypeptide chain and C349 of the second polypeptide chain.

In certain embodiments, a tetramer is provided, comprising a dimer of two DhCpmFc(+)(S354C)-GDF15(N3D):DhCpmFc(-)(Y349C) heterodimers in which the two first polypeptide chains of each heterodimer are linked via an interchain disulfide bond between their respective GDF15 regions.

More particularly, in a specific embodiment, the tetramer comprises:

(a) two DhCpmFc(+)(S354C) domains (one each heterodimer) comprising the sequence:

APELLGGPSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVD
GVEVHNAKTKPREEQYNSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPA
PIEKTISKAKGQPREPQVYTLPPCRKEMTKNQVSLTCLVKGFYPSDIAVE
WESNGQPENNYKTTPVCLKSDGSFFLYSKLTVDKSRWQQGNVVFSCSVMHE
ALHNHYTQKSLSLSPG (SEQ ID NO:85),

(b) two DhCpmFc(-)(Y349C) domains (one each heterodimer) comprising the sequence:

APELLGGPSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVD
 GVEVHNAKTKPREEQYNSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPA
 5 PIEKTISKAKGQPREPQVCTLPSSREEMTKNQVSLTCLVKGFYPSDIAVE
 WESNGQPENNYDTTPPVLDSDGSFFLYSDLTVDKSRWQQGNVVFSCSVMHE
 ALHNHYTQKSLSLSPG (SEQ ID NO:284),

and

(c) two GDF15(N3D) polypeptides (one each heterodimer) comprising the sequence
 10 of SEQ ID NO:52.

In a preferred embodiment, the first polypeptide chain comprises the amino acid sequence:

APELLGGPSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVD
 GVEVHNAKTKPREEQYNSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPA
 15 PIEKTISKAKGQPREPQVYTLPPCRKEMTKNQVSLTCLVKGFYPSDIAVE
 WESNGQPENNYKTTPPVLKSDGSFFLYSKLTVDKSRWQQGNVVFSCSVMHE
 ALHNHYTQKSLSLSPGARDGDHCPLGPGRCCRLHTVRASLEDLGWADWVL
 SPREVQVTMCIGACPSQFRAANMHAQIKTSLHRLKPDTVPAPCCVPASYN
 PMVLIQKTDGTGVSQTQYDDLAKDCHCI (SEQ ID NO:88),

20 which is encoded by the nucleic acid sequence:

gccccagagctgcttggtggaccatccgtgttcctgtttcctccaaagcc
 gaaggacaccctgatgatctcaagaactccggaagtgacttgcgtcgctg
 tggacgtgtcacatgaggatccagaggtcaagttcaattggtatgtggac
 ggagtggaagtgcataacgccaagaccaaaccggaagaacagtacaa
 25 tagcacctaccgctggtgagcgtccttactgtgctccaccaggactggc
 ttaatgggaaggaatacaagtgaaggtgtccaacaaggccctccccgct
 cccatcgaaaagaccatctcaaaggcaagggaaccaaagggaacctca
 agtgtacaccctgcctccgtgcaggaaggagatgaccaagaaccagggtca
 gcctgacttgctcgtgaagggtcttatccagcgatattgctgtggaa
 30 tgggagtgcaaatggccagcccagaataactacaaaactaccccaccgct
 gctgaaatctgatgggtccttcttcttactccaagctgaccgtggaca
 agagccgctggcaacaaggcaatgtctttagctgctcagtgatgcatgag
 gctctccataatcactacactcagaagtcactgtccctgtcacctggcgc
 gcgcgacggagaccactgtccgctcgggccccggcgcttgctgccgtctgc

acacgggtccgcgcgtcgctggaagacctgggctgggccgattgggtgctg
 tcgccacgggaggtgcaagtgacatgtgcacgcgtgccgagcca
 gttccggggcggcaaacatgcacgcgcagatcaagacgagcctgcaccgcc
 tgaagcccgacacggtgccagcgccctgctgcgtgcccgccagctacaat
 5 cccatggtgctcattcaaaagaccgacaccggggtgtcgctccagacct
 tgatgacttgttagccaaagactgccactgcata (SEQ ID NO:87).

In a preferred embodiment, the second polypeptide chain and comprises the amino acid sequence:

APELLGGPSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVD
 10 GVEVHNAKTKPREEQYNSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPA
 PIEKTISKAKGQPREPQVCTLPSSREEMTKNQVSLTCLVKGFYPSDIAVE
 WESNGQPENNYDTTPPVLDSDGSFFLYSDLTVDKSRWQQGNVFSQSVME
 ALHNHYTQKSLSLSPGK (SEQ ID NO:86),

which is encoded by the nucleic acid sequence:

gcacctgaactcctggggggaccgtcagtccttctcttccccccaaaacc
 caaggacaccctcatgatctcccgaccctgaggtcacatgcgtggtgg
 tggacgtgagccacgaagaccctgaggtcaagttcaactggtacgtggac
 ggcgtggaggtgcataatgccaaagacaaagccgcgggaggagcagtacaa
 cagcacgtaccgtgtggtcagcgtcctcaccgtcctgcaccaggactggc
 20 tgaatggcaaggagtacaagtgaaggtctccaacaaagccctccagcc
 cccatcgagaaaaccatctccaaagccaaagggcagccccgagaaccaca
 ggtgtgcaccctgcccccatcccgaggagatgaccaagaaccagggtca
 gcctgacctgcctggtcaaaggcttctatccagcgacatcgccgtggag
 tgggagagcaatgggcagccggagaacaactacgacaccacgcctccgt
 25 gctggactccgacggtccttcttctctatagcgacctcaccgtggaca
 agagcaggtggcagcaggggaacgtcttctcatgctccgtgatgcatgag
 gctctgcacaaccactacacgcagaagagcctctccctgtctccgggtaa
 a (SEQ ID NO:89).

As discussed above, a tetramer is provided comprising two polypeptide chains
 30 comprising the sequence of SEQ ID NO:88 and two polypeptide chains comprising the
 sequence of SEQ ID NO:86.

II.F.2 DhCpmFc(-)(Y349C)-GDF15:DhCpmFc(+)(S354C)

The designation “DhCpmFc(-)(Y349C)-GDF15:DhCpmFc(+)(S354C)” in the instant disclosure refers to a heterodimer comprising (i) a first polypeptide chain comprising a GDF15 polypeptide, the N-terminus of which is linked directly to the C-terminus of a DhCpmFc(-)(Y349C) domain, and (ii) a second polypeptide chain comprising a DhCpmFc(+)(S354C) domain. The cysteine clamp mutations allow the first and second polypeptide chains to be linked via an interchain disulfide bond between C349 of the first polypeptide chain and C354 of the second polypeptide chain.

In certain embodiments, a tetramer is provided, comprising a dimer of two DhCpmFc(-)(Y349C)-GDF15:DhCpmFc(+)(S354C) heterodimers in which the two first polypeptide chains of each heterodimer are linked via an interchain disulfide bond between their respective GDF15 regions.

More particularly, in a specific embodiment, the tetramer comprises:

(a) two DhCpmFc(+)(S354C) domains (one each heterodimer) comprising the sequence:

15 APELLGGPSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVD
GVEVHNAKTKPREEQYNSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPA
PIEKTISKAKGQPREPQVYTLPPCRKEMTKNQVSLTCLVKGFYPSDIAVE
WESNGQPENNYKTTPPVLKSDGSFFLYSKLTVDKSRWQQGNVFSQVMHE
ALHNHYTQKSLSLSPG (SEQ ID NO:285),

20 (b) two DhCpmFc(-)(Y349C) domains (one each heterodimer) comprising the sequence:

APELLGGPSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVD
GVEVHNAKTKPREEQYNSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPA
PIEKTISKAKGQPREPQVCTLPSSREEMTKNQVSLTCLVKGFYPSDIAVE
25 WESNGQPENNYDTTPPVLDSDGSFFLYSDLTVDKSRWQQGNVFSQVMHE
ALHNHYTQKSLSLSPG (SEQ ID NO:91),

and

(c) two GDF15 polypeptides (one each heterodimer) comprising the sequence of SEQ ID NO:12.

30 In a preferred embodiment, the first polypeptide chain comprises the amino acid sequence:

APELLGGPSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVD
GVEVHNAKTKPREEQYNSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPA
PIEKTISKAKGQPREPQVCTLPSSREEMTKNQVSLTCLVKGFYPSDIAVE

WESNGQPENNYDTPPVLDSDGSFFLYSDLTVDKSRWQQGNVFSVMSHE
 ALHNHYTQKSLSLSPGARNGDHCPLGPGRCCRLHTVRASLEDLGWADWVL
 SPREVQVTCIGACPSQFRAANMHAQIKTSLHRLKPDTPAPCCVPASYN
 PMVLIQKTDGTGVSLQTYDDLLAKDCHCI (SEQ ID NO:93),

5 which is encoded by the nucleic acid sequence:

gcacctgaactcctggggggaccgtcagtccttcctcttccccccaaaacc
 caaggacaccctcatgatctcccgaccctgaggtcacatgcgtggtgg
 tggacgtgagccacgaagaccctgaggtcaagttcaactggtacgtggac
 ggcgtggaggtgcataatgccaaagacaaagccgcgggaggagcagtacaa
 10 cagcacgtaccgtgtggtcagcgtcctcaccgtcctgcaccaggactggc
 tgaatggcaaggagtacaagtgaaggtctccaacaaagccctcccagcc
 cccatcgagaaaaccatctccaaagccaaagggcagccccgagaaccaca
 ggtgtgcaccctgcccccatcccgaggagatgaccaagaaccaggtca
 gcctgacctgcctggtcaaaggcttctatcccagcgacatcgccgtggag
 15 tgggagagcaatgggcagccggagaacaactacgacaccacgcctcccg
 gctggactccgacggctccttcttctctatagcgacctaccgtggaca
 agagcaggtggcagcaggggaacgtcttctcatgctccgtgatgcatgag
 gctctgcacaaccactacacgcagaagagcctctccctgtctccgggtgc
 ggcgaacggagaccactgtccgctcgggccccgggcgttgctgccgtctgc
 20 acacggtccgcgcgtcgctggaagacctgggctgggccgattgggtgctg
 tcgccacgggaggtgcaagtgacctgtgcatcggcgcgtgcccagacca
 gttccgggcggaacatgcacgcgcagatcaagacgagcctgcaccgcc
 tgaagcccgcacgggtgccagcgccctgctgcgtgcccgcagctacaat
 cccatggtgctcattcaaaagaccgacaccggggtgtcgctccagacct
 25 tgatgacttgtagccaaagactgccactgcata (SEQ ID NO:92).

In a preferred embodiment, the second polypeptide chain comprises the sequence:

APELLGGPSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVD
 GVEVHNAKTKPREEQYNSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPA
 PIEKTISKAKGQPREPQVYTLPPCRKEMTKNQVSLTCLVKGFYPSDIAVE
 30 WESNGQPENNYKTTPPVLSKSDGSFFLYSKLTVDKSRWQQGNVFSVMSHE
 ALHNHYTQKSLSLSPGK (SEQ ID NO:90),

which is encoded by the nucleic acid sequence:

gccccagagctgcttggtggaccatccgtgttcctgtttcctccaaagcc
 gaaggacaccctgatgatctcaagaactccggaagtgacttgctgcgtcg

tggacgtgtcacatgaggatccagaggtcaagttcaattggtatgtggac
 ggagtggaaagtgcataacgcccaagaccaaaccggaagaacagtacaa
 tagcacctaccgctggtgagcgtccttactgtgctccaccaggactggc
 ttaatgggaaggaataacaagtgttaaggtgtccaacaaggccctccccgt
 5 cccatcgaaaagaccatctcaaaggcaaaggggcaaccaagggaacctca
 agtgtacaccctgcctccgtgcaggaaggagatgaccaagaaccagggtca
 gcctgacttgtctcgtgaagggcttctatcccagcgatattgctgtggaa
 tgggagtcaaattggccagcccagagaataactacaaaactacccaccctgt
 gctgaaatctgatgggtccttcttcttcttactccaagctgaccgtggaca
 10 agagccgctggcaacaaggcaatgtcttttagctgctcagtcatgcatgag
 gctctccataatcactacactcagaagtcactgtccctgtctccgggtaa
 a (SEQ ID NO:94) .

As discussed above, a tetramer is provided comprising two polypeptide chains
 comprising the sequence of SEQ ID NO:93 and two polypeptide chains comprising the
 15 sequence of SEQ ID NO:90.

II.F.3 DhCpmFc(-)(Y349C)-GDF15(N3D):DhCpmFc(+)(S354C)

The designation “DhCpmFc(-)(Y349C)-GDF15(N3D):DhCpmFc(+)(S354C)” in the
 instant disclosure refers to heterodimer comprising (i) a first polypeptide chain comprising a
 20 GDF15(N3D) polypeptide, the N-terminus of which is linked directly to the C-terminus of a
 DhCpmFc(-)(Y349C) domain and (ii) a second polypeptide chain comprising a
 DhCpmFc(+)(S354C) domain. The cysteine clamp mutations allow the first and second
 polypeptide chains to be linked via an interchain disulfide bond between C349 of the first
 polypeptide and C354 of the second polypeptide.

25 In certain embodiments, a tetramer is provided, comprising a dimer of two
 DhCpmFc(-)(Y349C)-GDF15(N3D):DhCpmFc(+)(S354C) heterodimers in which the two
 first polypeptide chains of each heterodimer are linked via an interchain disulfide bond
 between their respective GDF15 regions. An N3D mutation in GDF15 sequence may be
 introduced, *e.g.*, to eliminate heterogeneity caused by N deamidation.

30 More particularly, in a specific embodiment, the tetramer comprises:

(a) two DhCpmFc(+)(S354C) domains (one each heterodimer) comprising the
 sequence of SEQ ID NO:285,

(b) two DhCpmFc(-)(Y349C) domains (one each heterodimer) comprising the
 sequence of SEQ ID NO:91, and

35 (c) two GDF15(N3D) polypeptides (one each heterodimer) comprising the sequence
 of SEQ ID NO:52.

In a preferred embodiment, the first polypeptide chain comprises the amino acid sequence:

APELLGGPSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVD
 GVEVHNAKTKPREEQYNSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPA
 5 PIEKTISKAKGQPREPQVCTLPPSREEMTKNQVSLTCLVKGFYPSDIAVE
 WESNGQPENNYDTPPVLDSDGSFFLYSDLTVDKSRWQQGNVFSCSVMHE
 ALHNHYTQKSLSLSPGARDGDHCPLGPGRCCRLHTVRASLEDLGWADWVL
 SPREVQVTMCIGACPSQFRAANMHAQIKTSLHRLKPDTPAPCCVPASYN
 PMVLIQKTDGTGVS LQTYDDLAKDCHCI (SEQ ID NO:96),

10 which is encoded by the nucleic acid sequence:

gcacctgaactcctggggggaccgtcagtccttcctcttccccccaaaacc
 caaggacaccctcatgatctcccgaccctgaggtcacatgcgtggtgg
 tggacgtgagccacgaagaccctgaggtcaagttcaactggtacgtggac
 ggcgtggaggtgcataatgccaaagacaaagccgcgggaggagcagtacaa
 15 cagcacgtaccgtgtggtcagcgtcctcaccgtcctgcaccaggactggc
 tgaatggcaaggagtacaagtgaaggtctccaacaaagccctcccagcc
 cccatcgagaaaaccatctccaaagccaaagggcagccccgagaaccaca
 ggtgtgcaccctgcccccatcccgaggagatgaccaagaaccaggtca
 gcctgacctgcctggtcaaaggcttctatcccagcgacatcgccgtggag
 20 tgggagagcaatgggcagccggagaacaactacgacaccacgcctcccgt
 gctggactccgacggtccttcttctctatagcgacctcaccgtggaca
 agagcaggtggcagcaggggaacgtcttctcatgctccgtgatgcatgag
 gctctgcacaaccactacacgcagaagagcctctccctgtctccgggtgc
 gcgcgacggagaccactgtccgctcgggcccggcggttgctgccgtctgc
 25 acacggtccgcgcgtcgctggaagacctgggctgggcccattgggtgctg
 tcgccacgggaggtgcaagtgaccatgtgcatcggcgcgtgcccgagcca
 gttccggggcggcaaacatgcacgcgcagatcaagacgagcctgcaccgcc
 tgaagcccgcacacggtgccagcgccctgctgcgtgcccgccagctacaat
 cccatgggtgctcattcaaaagaccgacaccggggtgtcgctccagaccta
 30 tgatgacttgtagccaaagactgccactgcata (SEQ ID NO:95).

In a preferred embodiment, the second polypeptide chain comprises the amino acid sequence of SEQ ID NO:90, which is encoded by the nucleic acid sequence of SEQ ID NO:94.

As discussed above, a tetramer is provided comprising two polypeptide chains comprising the sequence of SEQ ID NO:96 and two polypeptide chains comprising the sequence of SEQ ID NO:90.

5 *II.F.4 DhCpmFc(-)(Y349C)-GDF15(Ndel3):DhCpmFc(+)(S354C)*

The designation “DhCpmFc(-)(Y349C)-GDF15(Ndel3):DhCpmFc(+)(S354C)” in the instant disclosure refers to a heterodimer comprising (i) a first polypeptide chain comprising a GDF15(Ndel3) polypeptide, the N-terminus of which is linked directly to the C-terminus of a DhCpmFc(-)(Y349C) domain, and (ii) a second polypeptide chain comprising a
10 DhCpmFc(+)(S345C) domain. The cysteine clamp mutations allow the first and second polypeptide chains to be linked via an interchain disulfide bond between C349 of the first polypeptide chain and C354 of the second polypeptide chain.

In certain embodiments, a tetramer is provided, comprising a dimer of two DhCpmFc(-)(Y349C)-GDF15(Ndel3):DhCpmFc(+)(S354C) heterodimers in which the two
15 first polypeptide chains of each heterodimer are linked via an interchain disulfide bond between their respective GDF15 regions. Deletion of the N-terminal 3 amino acids (Ndel3) in GDF15 sequence may be introduced, *e.g.*, to eliminate heterogeneity caused by deamidation of asparagines or isomerization of aspartic acid.

More particularly, in a specific embodiment, the tetramer comprises:

20 (a) two DhCpmFc(+)(S354C) domains (one each heterodimer) of comprising the sequence of SEQ ID NO:285,

(b) two DhCpmFc(-)(Y349C) domains (one each heterodimer) comprising the sequence of SEQ ID NO:91, and

25 (c) two GDF15(Ndel3) polypeptides (one each heterodimer) comprising the sequence of SEQ ID NO:55.

In a preferred embodiment, the first polypeptide chain comprises the amino acid sequence:

30 APELLGGPSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVD
GVEVHNAKTKPREEQYNSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPA
PIEKTISKAKGQPREPQVCTLPSSREEMTKNQVSLTCLVKGFYPSDIAVE
WESNGQPENNYDTTPVLDSDGSFFLYSDLTVDKSRWQQGNVFSQSVME
ALHNHYTQKSLSLSPGGDHCPLGPGRCCRLHTVRASLEDLGWADWVLSR
EVQVTMCIGACPSQFRAANMHAQIKTSLHRLKPDTPAPCCVPASYNPMV
LIQKTDGTGVSLLQTYDDLLAKDCHCI (SEQ ID NO:98),

35 which is encoded by the nucleic acid sequence:

gcacctgaactcctggggggaccgtcagtccttcttccccccaaaacc
 caaggacaccctcatgatctcccgaccctgaggtcacatgcgtggtgg
 tggacgtgagccacgaagaccctgaggtcaagttcaactggtacgtggac
 ggcgtggaggtgcataatgccaaagacaaagccgcgggaggagcagtacaa
 5 cagcacgtaccgtgtggtcagcgtcctcaccgtcctgcaccaggactggc
 tgaatggcaaggagtacaagtgaaggtctccaacaaagccctcccagcc
 cccatcgagaaaaccatctccaaagccaaagggcagccccgagaaccaca
 ggtgtgcaccctgcccccatcccgaggagatgaccaagaaccaggtca
 gcctgacctgcctggtcaaaggcttctatcccagcgacatcgccgtggag
 10 tgggagagcaatgggcagccggagaacaactacgacaccacgcctcccgt
 gctggactccgacggctccttcttctctatagcgacctcaccgtggaca
 agagcaggtggcagcaggggaacgtcttctcatgctccgtgatgcatgag
 gctctgcacaaccactacacgcagaagagcctctccctgtctccgggtgg
 agaccactgtccgctcgggccccggcggttgcctgctctgcacacgggtcc
 15 gcgcgtcgctggaagacctgggctgggcccattgggtgctgtcgccacgg
 gaggtgcaagtgacctgtgcatcggcgcgtgcccagccagttccgggc
 ggcaaacatgcacgcgcagatcaagacgagcctgcaccgcctgaagcccg
 acacggtgccagcgccctgctgctgcccgcagctacaatcccatggtg
 ctcatcctaaaagaccgacacccgggtgtcgtccagacctatgatgactt
 20 gttagccaaagactgccactgcata (SEQ ID NO:97).

In a preferred embodiment, the second polypeptide chain comprises the amino acid sequence of SEQ ID NO:90, which is encoded by the nucleic acid sequence of SEQ ID NO:94.

As discussed above, a tetramer is provided comprising two polypeptide chains comprising the sequence of SEQ ID NO:98 and two polypeptide chains comprising the sequence of SEQ ID NO:90.

II.F.5 DhCpmFc(-)(Y349C)-G₄-GDF15(N3D):DhCpmFc(+)(S354C)

The designation “DhCpmFc(-)(Y349C)-G₄-GDF15(N3D):DhCpmFc(+)(S354C)” in the instant disclosure refers to a heterodimer comprising (i) a first polypeptide chain comprising a GDF15(N3D) polypeptide linked to a DhCpmFc(-)(Y349C) domain via a linker comprising the sequence of SEQ ID NO:58 that connects the N-terminus of the GDF15(N3D) polypeptide to the C-terminus of a DhCpmFc(-)(Y349C) domain and (ii) a second polypeptide chain comprising a DhCpmFc(+)(S354C) domain. The cysteine clamp mutations allow the first and second polypeptide chains to be linked via an interchain disulfide bond between C349 of the first polypeptide chain and C354 of the second polypeptide chain.

In certain embodiments, a tetramer is provided, comprising a dimer of two DhCpmFc(-)(Y349C)-G₄-GDF15(N3D):DhCpmFc(+)(S354C) heterodimers in which the two first polypeptide chains of each heterodimer are linked via an interchain disulfide bond between their respective GDF15 regions.

5 More particularly, in a specific embodiment, the tetramer comprises:

(a) two DhCpmFc(+)(S354C) domains (one each heterodimer) of comprising the sequence of SEQ ID NO:285,

(b) two DhCpmFc(-)(Y349C) domains (one each heterodimer) comprising the sequence of SEQ ID NO:91,

10 (c) two GDF15(N3D) polypeptides (one each heterodimer) comprising the sequence of SEQ ID NO:52, and

(d) two polypeptide linkers (one each heterodimer) comprising the sequence of SEQ ID NO:58 each linking the N -terminus of a GDF15(N3D) polypeptide to the C-terminus of a DhCpmFc(-)(Y349C) domain via a peptide bond.

15 In a preferred embodiment, the first polypeptide chain comprises the amino acid sequence (linker sequence double underlined):

APELLGGPSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVD
GVEVHNAKTKPREEQYNSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPA
PIEKTISKAKGQPREPQVCTLPSSREEMTKNQVSLTCLVKGFYPSDIAVE
20 WESNGQPENNYDTPPVLDSDGSFFLYSDLTVDKSRWQQGNVFSQSVME
ALHNHYTQKSLSLSPGGGGGARDGDHCPLGPGRCRLHTVRASLEDLGWA
DWVLSPREVQVTMCIGACPSQFRAANMHAQIKTSLHRLKPDTVPAPCCVP
ASYNPMVLIQKTDGTGVSLLQTYDDLLAKDCHCI (SEQ ID NO:100),

which is encoded by the nucleic acid sequence:

25 gcacctgaactcctggggggaccgtcagtccttcctcttccccccaaaacc
caaggacaccctcatgatctcccgaccctgaggtcacatgcgtggtgg
tggacgtgagccacgaagaccctgaggtcaagttcaactggtacgtggac
ggcgtggaggtgcataatgccaaagacaaagccgaggaggagcagtacaa
cagcacgtaccgtgtggtcagcgtcctcaccgtcctgcaccaggactggc
30 tgaatggcaaggagtacaagtgaaggtctccaacaaagccctcccagcc
cccatcgagaaaaccatctccaaagccaaagggcagccccgagaaccaca
ggtgtgcaccctgcccccatcccgaggagatgaccaagaaccaggtca
gcctgacctgcctggtcaaaggcttctatcccagcgacatcgccgtggag
tgggagagcaatgggcagccggagaacaactacgacaccacgcctcccg
35 gctggactccgacggctccttcttctctatagcgacctcaccgtggaca

agagcaggtggcagcaggggaacgtcttctcatgctccgtgatgcatgag
gctctgcacaaccactacacgcagaagagcctctccctgtctccgggtgg
tggaggtggtgcgcgcgacggagaccactgtccgctcgggcccgggcggt
gctgccgtctgcacacgggtccgcgcgtcgctggaagacctgggctgggccc
5 gattgggtgctgtcgccacgggaggtgcaagtgacctgtgcatcgggcgc
gtgcccagaccagttccggggcggaacatgcacgcgcagatcaagacga
gcctgcaccgcctgaagcccgacacgggtgccagcgccctgctgcgtgccc
gccagctacaatcccatggtgctcattcaaaagaccgacaccgggggtgtc
gctccagacctatgatgacttgtagccaaagactgccactgcata (SEQ ID NO:99) .

10 In a preferred embodiment, the second polypeptide chain comprises the amino acid sequence of SEQ ID NO:90, which is encoded by the nucleic acid sequence of SEQ ID NO:94.

As discussed above, in a specific embodiment, a heterotetramer is provided comprising two monomers having the sequence of SEQ ID NO:100 and two monomers
15 having the sequence of SEQ ID NO:90.

II.F.6 DhCpmFc(-)(Y349C)-(G₄S)₂-GDF15(N3D):DhCpmFc(+)(S354C)

The designation “DhCpmFc(-)(Y349C)-(G₄S)₂-GDF15(N3D):DhCpmFc(+)(S354C)” in the instant disclosure refers to a heterodimer comprising (i) a first polypeptide chain
20 comprising a GDF15(N3D) polypeptide linked to a DhCpmFc(-)(Y349C) domain via a linker comprising the sequence of SEQ ID NO:64 that connects the N-terminus of the GDF15(N3D) polypeptide to the C-terminus of a DhCpmFc(-)(Y349C) domain, and (ii) a second polypeptide chain comprising a DhCpmFc(+)(S354C) domain. The cysteine clamp mutations allow the first and second polypeptide chains to be linked via an interchain disulfide bond
25 between C349 of the first polypeptide chain and C354 of the second polypeptide chain.

In certain embodiments, a tetramer is provided, comprising a dimer of two DhCpmFc(-)(Y349C)-(G₄S)₂-GDF15(N3D):DhCpmFc(+)(S354C) heterodimers in which the two first polypeptide chains of each heterodimer are linked via an interchain disulfide bond between their respective GDF15 regions.

30 More particularly, in a specific embodiment, the tetramer:

(a) two DhCpmFc(+)(S354C) domains (one each heterodimer) of comprising the sequence of SEQ ID NO:285,

(b) two DhCpmFc(-)(Y349C) domains (one each heterodimer) comprising the sequence of SEQ ID NO:91,

35 (c) two GDF15(N3D) polypeptides (one each heterodimer) comprising the sequence of SEQ ID NO:52, and

(d) two polypeptide linkers (one each heterodimer) comprising the sequence of SEQ ID NO:64 each linking the N-terminus of a GDF15(N3D) polypeptide to the C-terminus of a DhCpmFc(-)(Y349C) domain via a peptide bond.

In a preferred embodiment, the first polypeptide comprises the amino acid sequence
5 (linker sequence double underlined):

APELLGGPSVFLFPPKPKDTLMI SRTPEVTCVVVDVSHEDPEVKFNWYVD
GVEVHNAKTKPREEQYNSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPA
PIEKTISKAKGQPREPQVCTLPSSREEMTKNQVSLTCLVKGFYPSDIAVE
WESNGQPENNYDTPPVLDSDGSFFLYSDLTVDKSRWQQGNVFCFSVMHE
10 ALHNHYTQKSLSLSPGGGGSGGGGSARDGDHCPLGPGRCCRLHTVRASL
EDLGWADWVLSPREVQVTMCIGACPSQFRAANMHAQIKTSLHRLKPDTVP
APCCVPASYNPMVLIQKTDGTGVSLSQTYDDLAKDCHCI (SEQ ID NO:102),

which is encoded by the nucleic acid sequence:

gcacctgaactcctggggggaccgtcagtccttcctcttccccccaaaacc
15 caaggacaccctcatgatctcccgaccctgaggtcacatgcgtggtgg
tggacgtgagccacgaagaccctgaggtcaagttcaactggtacgtggac
ggcgtggaggtgcataatgccaagacaaagccgcgggaggagcagtacaa
cagcacgtaccgtgtggtcagcgtcctcaccgtcctgcaccaggactggc
tgaatggcaaggagtacaagtgaaggtctccaacaaagccctcccagcc
20 cccatcgagaaaaccatctccaaagccaaagggcagccccgagaaccaca
ggtgtgcaccctgcccccatcccgaggagatgaccaagaaccagggtca
gcctgacctgcctggtcaaaggcttctatcccagcgacatcgccgtggag
tgggagagcaatgggcagccggagaacaactacgacaccacgcctcccgt
gctggactccgacggctccttcttctctatagcgacctcaccgtggaca
25 agagcaggtggcagcaggggaacgtcttctcatgctccgtgatgcatgag
gctctgcacaaccactacacgcagaagagcctctccctgtctccgggtgg
aggtggtggatccggaggcgggtggaagcgcgcgcgacggagaccactgtc
cgctcggggcccggttgctgccgtctgcacacggtccgcgcgtcgctg
gaagacctgggctgggctgattgggtgctgtcgccacgggaggtgcaagt
30 gaccatgtgcatcggcgcgtgcccagagccagttccgggaggcaaacatgc
acgcgcagatcaagacgagcctgcaccgcctgaagcccgcacgggtgcc
gcgcctgctgcgtgcccgcagctacaatcccatggtgctcattcaaaa
gaccgacaccggggtgtcgctccagacctatgatgacttgtagccaaag
actgccactgcata (SEQ ID NO:101).

In a preferred embodiment, the second polypeptide chain comprises the amino acid sequence of SEQ ID NO:90, which is encoded by the nucleic acid sequence of SEQ ID NO:94.

- As discussed above, in a specific embodiment, a heterotetramer is provided
- 5 comprising two monomers having the sequence of SEQ ID NO:102 and two monomers having the sequence of SEQ ID NO:90.

II.F.7 DhCpmFc(-)(Y349C)-(G₄Q)₂-GDF15(N3D):DhCpmFc(+)(S354C)

- The designation “DhCpmFc(-)(Y349C)-(G₄Q)₂-GDF15(N3D):DhCpmFc(+)(S354C)”
- 10 in the instant disclosure refers to a heterodimer comprising (i) a first polypeptide chain comprising a GDF15(N3D) polypeptide linked to a DhCpmFc(-)(Y349C) domain via a linker comprising the sequence of SEQ ID NO:78 that connects the N-terminus of the GDF15(N3D) polypeptide to the C-terminus of the DhCpmFc(-)(Y349C) domain, and (ii) a second polypeptide chain comprising a DhCpmFc(+)(S354C) domain. The cysteine clamp mutations
- 15 allow the first and second polypeptide chains to be linked via an interchain disulfide bond between C349 of the first polypeptide chain and C354 of the second polypeptide chain.

- In certain embodiments, a tetramer is provided, comprising a dimer of two DhCpmFc(-)(Y349C)-(G₄Q)₂-GDF15(N3D):DhCpmFc(+)(S354C) heterodimers in which the two first polypeptide chains of each heterodimer are linked via an interchain disulfide bond
- 20 between their respective GDF15 regions.

More particularly, in a specific embodiment, the tetramer comprises:

- (a) two DhCpmFc(+)(S354C) domains (one each heterodimer) of comprising the sequence of SEQ ID NO:285,
- (b) two DhCpmFc(-)(Y349C) domains (one each heterodimer) comprising the
- 25 sequence of SEQ ID NO:91,
- (c) two GDF15(N3D) polypeptides (one each heterodimer) comprising the sequence of SEQ ID NO:52, and
- (d) two polypeptide linkers (one each heterodimer) comprising the sequence of SEQ ID NO:78 each linking the N-terminus of a GDF15(N3D) polypeptide to the C-terminus of a
- 30 DhCpmFc(-)(Y349C) domain via a peptide bond.

In a preferred embodiment, the first polypeptide comprises the amino acid sequence (linker sequence double underlined):

- APELLGGPSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVD
- GVEVHNAKTKPREEQYNSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPA
- 35 PIEKTISKAKGQPREPQVCTLPSSREEMTKNQVSLTCLVKGFYPSDIAVE
- WESNGQPENNYDTTPPVLDSDGSFFLYSDLTVDKSRWQQGNVVFSCSVME

ALHNHYTQKSLSLSPGGGGGQGGGGQARDGDHCPLGPGRCCRLHTVRASL
EDLGWADWVLSPREVQVTMCIGACPSQFRAANMHAQIKTSLHRLKPDTVP
APCCVPASYNPMVLIQKTDGTGVSLLQTYDDLLAKDCHCI (SEQ ID NO:104),

which is encoded by the nucleic acid sequence:

5 gcacctgaactcctggggggaccgtcagtccttcttccccccaaaacc
caaggacaccctcatgatctcccggaaccctgaggtcacatgcgtggtgg
tggacgtgagccacgaagaccctgaggtcaagttcaactggtacgtggac
ggcgtggaggtgcataatgccaaagacaaagccgcgggaggagcagtacaa
cagcacgtaccgtgtggtcagcgtcctcaccgtcctgcaccaggactggc
10 tgaatggcaaggagtacaagtgaaggtctccaacaaagccctcccagcc
cccatcgagaaaaccatctccaaagccaaagggcagccccgagaaccaca
ggtgtgcaccctgcccccatcccgaggagatgaccaagaaccaggtca
gcctgacctgcctggtcaaaggcttctatcccagcgacatcgccgtggag
tgggagagcaatgggcagccggagaacaactacgacaccacgcctcccg
15 gctggactccgacggctccttcttctctatagcgacctcaccgtggaca
agagcaggtggcagcaggggaacgtcttctcatgctccgtgatgcatgag
gctctgcacaaccactacacgcagaagagcctctccctgtctccgggtgg
aggtggtggacagggaggcgggtggacagggcgcgcgacggagaccactgtc
cgctcggggcccgggcggttgctgccgtctgcacacgggtccgcgcgtcgctg
20 gaagacctgggctgggcccattgggtgctgtcgccacgggaggtgcaagt
gaccatgtgcatcggcgcgtgcccagagccagttccgggaggcaaacatgc
acgcgcagatcaagacgagcctgcaccgcctgaagcccgacacgggtgcc
gcgccctgctgctgcccgcagctacaatcccatggtgctcattcaaaa
gaccgacaccggggtgtcgctccagacctatgatgacttgtagccaaag
25 actgccactgcata (SEQ ID NO:103).

In a preferred embodiment, the second polypeptide chain comprises the amino acid sequence of SEQ ID NO:90, which is encoded by the nucleic acid sequence of SEQ ID NO:94.

As discussed above, in a specific embodiment, a heterotetramer is provided
30 comprising two monomers having the sequence of SEQ ID NO:104 and two monomers having the sequence of SEQ ID NO:90.

II.F.8. DhCpmFc(-)(L351C)-(G₄S)₂-GDF15:DhCpmFc(+)(L351C)

The designation “DhCpmFc(-)(L351C)-(G₄S)₂-GDF15:DhCpmFc(+)(L351C)” in the
35 instant disclosure refers to a heterodimer comprising (i) a first polypeptide chain comprising a

GDF15 polypeptide linked to a DhCpmFc(-)(L351C) domain via a linker comprising the sequence of SEQ ID NO:64 that connects the N-terminus of the GDF15 polypeptide to the C-terminus of the DhCpmFc(-)(L351C) domain, and (ii) a second polypeptide chain comprising a DhCpmFc(+)(L351C) domain. The cysteine clamp mutations allow the first and second polypeptide chains to be linked via an interchain disulfide bond between C351 of the first polypeptide chain and C351 of the second polypeptide chain.

In certain embodiments, a tetramer is provided, comprising a dimer of two DhCpmFc(-)(L351C)-(G₄S)₂-GDF15:DhCpmFc(+)(L351C) heterodimers in which the two first polypeptide chains of each heterodimer are linked via an interchain disulfide bond between their respective GDF15 regions.

More particularly, in a specific embodiment, the tetramer comprises:

(a) two DhCpmFc(+)(L351C) domains (one each heterodimer) comprising the sequence:

APELLGGPSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVD
 GVEVHNAKTKPREEQYNSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPA
 PIEKTISKAKGQPREPQVYTCPPSRKEMTKNQVSLTCLVKGFYPSDIAVE
 WESNGQPENNYKTTPPVLKSDGSFFLYSKLTVDKSRWQQGNVFSQVMHE
 ALHNHYTQKSLSLSPG (SEQ ID NO:286),

(b) two DhCpmFc(-)(L351C) domains (one each heterodimer) comprising the sequence:

APELLGGPSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVD
 GVEVHNAKTKPREEQYNSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPA
 PIEKTISKAKGQPREPQVYTCPPSREEMTKNQVSLTCLVKGFYPSDIAVE
 WESNGQPENNYDTTPPVLDSDGSFFLYSDLTVDKSRWQQGNVFSQVMHE
 ALHNHYTQKSLSLSPG (SEQ ID NO:106),

(c) two GDF15 polypeptide chains (one each heterodimer) comprising the sequence of SEQ ID NO:12, and

(d) two polypeptide linkers (one each heterodimer) comprising the sequence of SEQ ID NO:64 each linking the N-terminus of a GDF15 polypeptide to the C-terminus of a DhCpmFc(-)(L351C) domain via a peptide bond.

In a preferred embodiment, the first polypeptide chain comprises the amino acid sequence (linker sequence double underlined):

APELLGGPSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVD
 GVEVHNAKTKPREEQYNSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPA

PIEKTISKAKGQPREPQVYTCPPSREEMTKNQVSLTCLVKGFYPSDIAVE
 WESNGQPENNYDTPPVLDSDGSFFLYSDLTVDKSRWQQGNVFSQSVHME
 ALHNHYTQKSLSLSPGGGGGSGGGGSARNGDHCPLGPGRCCRLHTVRASL
 EDLGWADWVLSPREVQVTMCIGACPSQFRAANMHAQIKTSLHRLKPDTPV
 5 APCCVPASYNPMVLIQKTDGTGVSLLQTYDDLLAKDCHCI (SEQ ID NO:108),

which is encoded by the nucleic acid sequence:

gcgcgcgaactgctggcgcccgagcgtgtttctgtttccgcgaaacc
 gaaagataccctgatgattagccgcaccccggaagtacctgctggtgg
 tggatgtgagccatgaagatccggaagtgaatttaactggtatgtggat
 10 ggctggaagtgcataacgcgaaaaccaaaccgcgcgaagaacagtataa
 cagcacctatcgctggtgagcgtgctgaccgtgctgcatcaggattggc
 tgaacggcaaagaatataaatgcaaagtgagcaaaaagcgtgccggcg
 ccgattgaaaaaacattagcaaagcgaaaggccagccgcgcgaaccgca
 ggtgtatacctgcccgcgagccgcgaagaaatgacaaaaaccaggtga
 15 gcctgacctgcctggtgaaaggcttttatccgagcgatattgcggtggaa
 tgggaaagcaacggccagccggaacaactatgataccaccccgccggt
 gctggatagcgatggcagctttttctgtatagcgatctgaccgtggata
 aaagccgctggcagcagggcaacgtgttttagctgcagcgtgatgcatgaa
 gcgctgcataaccattataccagaaaagcctgagcctgagccggggcg
 20 cgggcgccgagcggcgccggcgccgagcgcgcgcaacggcgatcattgcc
 cgctgggcccggccgctgctgcccgcctgcataccgtgcgcgcgagcctg
 gaagatctgggctgggaggattgggtgctgagcccgcgcaagtgcaggt
 gaccatgtgcattggcgcgtgcccagagccagtttcgcgcggcgaaatgc
 atgcgcagattaaaaccagcctgcctgcctgaaaccggataccgtgccg
 25 gcgcctgctgctgctgccggcgagctataaccgatggtgctgattcagaa
 aaccgataccggcgtgagcctgcagacctatgatgatctgctggcgaaag
 attgccattgcatt (SEQ ID NO:107).

In a preferred embodiment, the second polypeptide chain and comprises the amino acid sequence:

30 APELLGGPSVFLFPPKPKDTLMI SRTPEVTCVVVDVSHEDPEVKFNWYVD
 GVEVHNAKTKPREEQYNSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPA
 PIEKTISKAKGQPREPQVYTCPPSRKEMTKNQVSLTCLVKGFYPSDIAVE
 WESNGQPENNYKTTPPVLKSDGSFFLYSKLTVDKSRWQQGNVFSQSVHME
 ALHNHYTQKSLSLSPGK (SEQ ID NO:105),

which is encoded by the nucleic acid sequence:

```

gcacctgaactcctggggggaccgtcagtcttctcttccccccaaaacc
caaggacaccctcatgatctcccgaccctgaggtcacatgcgtggtgg
tggacgtgagccacgaagaccctgaggtcaagttcaactggtacgtggac
5  ggcgtggaggtgcataatgccaagacaaagccgcgggaggagcagtacaa
cagcacgtaccgtgtggtcagcgtcctcaccgtcctgcaccaggactggc
tgaatggcaaggagtacaagtgaagggtctccaacaaagccctcccagcc
cccatcgagaaaaccatctccaaagccaaagggcagccccgagaaccaca
ggtgtacacctgtcccccatcccgaggagatgaccaagaaccagggtca
10  gcctgacctgcctggtcaaaggcttctatcccagcgacatcgccgtggag
tgggagagcaatgggcagccggagaacaactacgacaccacgcctcccg
gctggactccgacggctccttcttctctatagcgacctcaccgtggaca
agagcaggtggcagcaggggaacgtcttctcatgctccgtgatgcatgag
gctctgcacaaccactacacgcagaagagcctctccctgtctccgggtaa
15  a (SEQ ID NO:109) .

```

As discussed above, in a specific embodiment, a heterotetramer is provided comprising two monomers having the sequence of SEQ ID NO:108 and two monomers having the sequence of SEQ ID NO:105.

20 II.G. HSA

The designations “HSA” or “human serum albumin” in the instant disclosure refer to a fusion protein comprising a GDF15 region linked, directly or via a polypeptide linker, to a human serum albumin (HSA) polypeptide. In some embodiments, the fusion protein comprises two or more HSA polypeptides.

25 Typically, the N-terminus of the GDF15 region is linked, directly or via a polypeptide linker, to the C-terminus of the HSA polypeptide. However, in some embodiments, the N-terminus of the HSA polypeptide is linked, directly or via a polypeptide linker, to the C-terminus of the GDF15 region.

In certain embodiments, a homodimer is provided comprising two such fusion
30 proteins linked via an interchain disulfide bond between their respective GDF15 regions. *See* Figure 6 for a graphic depiction of an embodiment of such a homodimer. Alternatively, a heterodimer is provided comprising one such fusion protein and a GDF15 polypeptide or GDF mutant polypeptide, linked via an interchain disulfide bond between the GDF15 region of the fusion protein and the GDF15 polypeptide or mutant polypeptide.

35

II.G.1 HSA-(G₄S)₄-GDF15:GDF15 heterodimer

- The designation “HSA-(G₄S)₄-GDF15:GDF15” in the instant disclosure refers to a heterodimer comprising (i) a first polypeptide chain comprising a GDF15 polypeptide linked to an HSA polypeptide via a linker comprising the sequence of SEQ ID NO:18 that connects the N-terminus of the GDF15 polypeptide to the C-terminus of the HSA polypeptide, and (ii)
- 5 a second polypeptide chain comprising a GDF15 polypeptide.

Typically, the first and second polypeptide chains are linked via an interchain disulfide bond between their respective GDF15 regions.

More particularly, in a specific embodiment, the heterodimer comprises:

(a) one HSA polypeptide (first monomer) comprising the sequence:

10 DAHKSEVAHREFKDLGEENFKALVLI~~AF~~AQYLQQCPFEDHVKLVNEVTEFA
KTCVADESAENCDKSLHTLFGDKLCTVATLRETYGEMADCCAKQEPERNE
CFLQHKDDNPNL~~PRL~~VLRPEVDVMCTAFHDNEETFLKKYLYEIARRHPYFY
APELLFFAKRYKAAFTECCQAADKAACLLPKLDEL~~RDE~~GKASSAKQRLKC
ASLQKFGERAFKAWAVARLSQRFPKAEFAEVSKLVTDLTKVHTECCHGDL
15 LECADDRADLAKYICENQDSISSKLKECCEKPLEKSHCIAEVENDEMPA
DLPSLAADFVESKDVCKNYAEAKDVFLGMFLYEYARRHPDYSVVL~~LL~~RLA
KTYETTTLEKCCAAADPHECYAKVFDEFKPLVEEPQNLIKQNC~~EL~~FEQLGE
YKFQNALLVRYTKKVPQVSTPTLVEVSRNLGKVGSKCKHPEAKRMPCAE
DYL~~SV~~VLNQLC~~VL~~HEKTPVSDRVTKCTESLVNRRPCFSALEVD~~ET~~YVPK
20 EFNAETFTFHADICTLSEKERQIKKQTALVELVKHKPKATKEQLKAVMDD
FAAFVEKCKKADDKETCFAEEGKKLVAA~~SQA~~ALGL (SEQ ID NO:110),

and

(b) two GDF15 polypeptides (one each monomer) comprising the sequence of SEQ ID NO:12, and

- 25 (c) one polypeptide linker (first monomer) comprising the sequence of SEQ ID NO:18 linking the N-terminus of a GDF15 polypeptide to the C-terminus of the HSA polypeptide via a peptide bond.

In a preferred embodiment, the first polypeptide comprises the amino acid sequence (linker double underlined):

30 DAHKSEVAHREFKDLGEENFKALVLI~~AF~~AQYLQQCPFEDHVKLVNEVTEFA
KTCVADESAENCDKSLHTLFGDKLCTVATLRETYGEMADCCAKQEPERNE
CFLQHKDDNPNL~~PRL~~VLRPEVDVMCTAFHDNEETFLKKYLYEIARRHPYFY
APELLFFAKRYKAAFTECCQAADKAACLLPKLDEL~~RDE~~GKASSAKQRLKC
ASLQKFGERAFKAWAVARLSQRFPKAEFAEVSKLVTDLTKVHTECCHGDL
35 LECADDRADLAKYICENQDSISSKLKECCEKPLEKSHCIAEVENDEMPA
DLPSLAADFVESKDVCKNYAEAKDVFLGMFLYEYARRHPDYSVVL~~LL~~RLA

KTYETTLEKCCAAADPHECYAKVFDEFKPLVEEPQNLIKQNCSELFELGE
 YKFQNALLVRYTKKVPQVSTPTLVEVSRNLGKVGSKCKHPEAKRMPCAE
 DYLSVVLNQLCVLHEKTPVSDRVTKCTESLVNRRPCFSALEVDETYVPK
 EFNAETFTFHADICTLSEKERQIKKQTALVELVKHKPKATKEQLKAVMDD
 5 FAAFVEKCKKADDKETCFAEEGKKLVAASQAALGLGGGSGGGSGGGGS
GGGGSARNGDHCPLGPGRCCRLHTVRASLEDLGWADWVLSPREVQVTMCI
 GACPSQFRAANMHAQIKTSLHRLKPDTPAPCCVPASYNPMVLIQKTDTG
 VSLQTYDDLAKDCHCI (SEQ ID NO:112),

which is encoded by the nucleic acid sequence:

10 gatgcacacaagagtgaggttgctcatcgatttaaagatttgggagaaga
 aaatttcaaagccttggtggtgattgccttgctcagtatcttcagcagt
 gtccatttgaagatcatgtaaaattagtgaatgaagtaactgaatttgca
 aaaacatgtgttgctgatgagtcagctgaaaattgtgacaaatcacttca
 taccctttttggagacaaattatgcacagttgcaactcttcgtgaaacct
 15 atggtgaaatggctgactgctgtgcaaaacaagaacctgagagaaatgaa
 tgcttcttgcaacacaaagatgacaacccaaacctccccgattggtgag
 accagaggttgatgtgatgtgcactgcttttcatgacaatgaagagacat
 ttttgaaaaaataacttatatgaaattgccagaagacatccttacttttat
 gccccggaactccttttcttgctaaaaggtataaagctgcttttacaga
 20 atgttgccaagctgctgataaagctgcctgcctgttgccaaagctcgatg
 aacttcgggatgaagggaaggcttcgtctgccaacagagactcaagtgt
 gccagtctccaaaaatttggagaaagagcttcaaagcatgggcagtagc
 tcgcctgagccagagatttccaaagctgagtttgagaagtttccaagt
 tagtgacagatcttaccaaagtcacacggaatgctgccatggagatctg
 25 cttgaatgtgctgatgacagggcggaaccttgccaagtatatctgtgaaaa
 tcaagattcgatctccagtaaaactgaaggaatgctgtgaaaaacctctgt
 tggaaaaatcccactgcattgccgaagtggaaaaatgatgagatgcctgct
 gacttgccttcattagctgctgattttgttgaaagtaaggatgtttgcaa
 aaactatgctgaggcaaaggatgtcttccctgggcatgtttttgtatgaat
 30 atgcaagaaggcatcctgattactctgtcgtgctgctgctgagacttgcc
 aagacatatgaaaccactctagagaagtgctgtgccgctgcagatcctca
 tgaatgctatgccaaagtgttcgatgaatttaaacctcttggtggaagagc
 ctcagaatttaatacaacaaaattgtgagctttttgagcagcttgagagag
 taaaaattccagaatgcgctattagttcgttacaccaagaaagtacccca
 35 agtgtcaactccaactcttgtagaggtctcaagaaacctagggaaagtgg
 gcagcaaatgttgtaaacatcctgaagcaaaaagaatgcctgtgcagaa

gactatctatccgtggctcctgaaccagttatgtgtgttgcattgagaaaac
 gccagtaagtgcagagtcaccaaagctgcacagaatccttggatgaaca
 ggcgaccatgcttttcagctctggaagtcgatgaaacatacgttccaaa
 gagtttaagtctgaaacattcaccttccatgcagatatatgcacactttc
 5 tgagaaggagagacaaatcaagaaacaaactgcacttgttgagctcgtga
 aacacaagcccaaggcaacaaaagagcaactgaaagctgttatggatgat
 ttctgcagctttttagagaagtgcagcaaggctgacgataaggagacctg
 ctttgccgaggagggtaaaaaacttgttgccggcagtcaggccgccttag
 gcttaggaggtggtggatccggaggcggtggaagcggaggtggtggatct
 10 ggaggcggtggaagcgcgcgcaacggagaccactgtccgctcgggcccgg
 gcgttgctgccgtctgcacacggctccgcgcgtcgtggaagacctgggct
 gggccgattgggtgctgctgccacgggaggtgcaagtaccatgtgcac
 ggcgcgtgcccagaccagttccggggcgcaaacatgcacgcgcagatcaa
 gacgagcctgcaccgcctgaagcccgacacgggtgccagcgcctgctgcg
 15 tgcccgccagctacaatcccatggtgctcattcaaaagaccgacaccggg
 gtgctcgtccagacctatgatgacttgttagccaaagactgccactgcat
 atga (SEQ ID NO:111).

In a preferred embodiment, the second polypeptide chain comprises the amino acid
 sequence of SEQ ID NO:12, which is encoded by the nucleic acid sequence of SEQ ID
 20 NO:11.

As discussed above, a heterodimer is provided comprising a first polypeptide chain
 having the sequence of SEQ ID NO:112 and a second polypeptide chain having the sequence
 of SEQ ID NO:12.

25 II.G.2 HSA-(G₄S)₄-GDF15

The designation “HSA-(G₄S)₄-GDF15” in the instant disclosure refers to a fusion
 protein comprising a GDF15 polypeptide linked to an HSA polypeptide via a linker
 comprising the sequence of SEQ ID NO:18 that connects the N-terminus of the GDF15
 polypeptide to the C-terminus of the HSA polypeptide.

30 In certain embodiments, a homodimer is provided comprising two HSA-(G₄S)₄-
 GDF15 fusion proteins, linked via an interchain disulfide bond between their respective
 GDF15 regions.

More particularly, in a specific embodiment, the homodimer comprises:

(a) two HSA polypeptides (one each monomer) comprising the sequence of SEQ ID
 35 NO:110;

(b) two GDF15 polypeptides (one each monomer) comprising the sequence of SEQ ID NO:12, and

(c) two polypeptide linkers (one each monomer) comprising the sequence of SEQ ID NO:18 each linking the N-terminus of a GDF15 polypeptide to the C-terminus of an HSA polypeptide via a peptide bond.

In a preferred embodiment, the fusion protein comprises the amino acid sequence of SEQ ID NO:112, which is encoded by the nucleic acid sequence of SEQ ID NO:111.

As discussed above, in a specific embodiment, a homodimer is provided comprising two fusion proteins having the sequence of SEQ ID NO:112.

II.G.3 HSA-GSPAPAPGS-GDF15

The designation “HSA-(GSPAPAPGS)-GDF15” in the instant disclosure refers to a fusion protein comprising a GDF15 polypeptide linked to an HSA polypeptide via a linker comprising the sequence of SEQ ID NO:113 that connects the N-terminus of the GDF15 polypeptide to the C-terminus of the HSA polypeptide.

In certain embodiments, a homodimer is provided comprising two HSA-(GSPAPAPGS)-GDF15 fusion proteins, linked via an interchain disulfide bond between their respective GDF15 regions.

More particularly, in a specific embodiment, the homodimer comprises:

(a) two HSA polypeptides (one each monomer) comprising the sequence of SEQ ID NO:110;

(b) two GDF15 polypeptides (one each monomer) comprising the sequence of SEQ ID NO:12; and

(c) two polypeptide linkers (one each monomer) comprising the sequence:

GSPAPAPGS (SEQ ID NO:113)

each linking the N-terminus of the GDF-15 polypeptide to the C-terminus of the HSA polypeptide via a peptide bond.

In a preferred embodiment, the fusion protein comprises the amino acid sequence (linker double underlined):

DAHKSEVAHREFKDLGEENFKALVLI~~AFAYLQQCPFEDHVKLVNEVTEFA~~
 KTCVADESAENCDKSLHTLFGDKLCTVATLRETYGEMADCCAKQEPERNE
 CFLQHKDDNPNLPRLVRPEVDVMCTAFHDNEETFLKKYLYEIARRHPYFY
 APELLFFAKRYKAAFTECCQAADKAACLLPKLDEL~~RDEGKASSAKQRLKC~~
 ASLQKFGERAFKAWAVARLSQRFPKAEFAEVSKLVTDLTKVHTECCHGDL
 LECADDRADLAKYICENQDSISSKLKECCEKPLEKSHCIAEVENDEMPA
 DLPSLAADFVESKDVCKNYAEAKDVFLGMFLY~~EYARRHPDYSVVL~~LLRLA

KTYETTTLEKCCAAADPHECYAKVFDEFKPLVEEPQNLIKQNCSELFQELGE
 YKFQNALLLVRYTKKVPQVSTPTLVEVSRNLGKVGSKCKHPEAKRMPCAE
 DYLSVVLNQLCVLHEKTPVSDRVTKCTESLVNRRPCFSALEVDETYVPK
 EFNAETFTFHADICTLSEKERQIKKQTALVELVKHKPKATKEQLKAVMDD
 5 FAAFVEKCKKADDKETCFAEEGKKLVAASQAALGLGSPAPAPGSARNGDH
 CPLGPGRCCRLHTVRASLEDLGWADWVLSPREVQVTCIGACPSQFRAAN
 MHAQIKTSLHRLKPDTPAPCCVPASYNPMVLIQKTDGTGVSLQTYDDLLA
 KDCHCI (SEQ ID NO:115),

which is encoded by the nucleic acid sequence:

10 gatgcacacaagagtgaggttgctcatcgatttaaagatttgggagaaga
 aaatttcaaagccttggtggtgattgccttgcctcagtatcttcagcagt
 gtccatttgaagatcatgtaaaattagtgaatgaagtaactgaatttgca
 aaaacatgtgttgctgatgagtcagctgaaaattgtgacaaatcacttca
 tacccttttggagacaaattatgcacagttgcaactcttcgtgaaacct
 15 atggtgaaatggctgactgctgtgcaaaacaagaacctgagagaaatgaa
 tgcttcttgcaacacaaagatgacaacccaaacctccccgattggtgag
 accagaggttgatgtgatgtgcactgcttttcatgacaatgaagagacat
 ttttgaaaaaatacttatatgaaattgccagaagacatccttacttttat
 gccccggaactccttttcttgctaaaaggtataaagctgcttttacaga
 20 atgttgccaagctgctgataaagctgcctgcctgttgccaaagctcgatg
 aacttcgggatgaagggaaggcttcgtctgccaacagagactcaagtgt
 gccagtctccaaaaatttggagaaagagcttcaaagcatgggcagtagc
 tcgcctgagccagagatttccaaagctgagtttgagaagtttccaagt
 tagtgacagatcttaccaaagtccacacggaatgctgccatggagatctg
 25 cttgaatgtgctgatgacagggcggaaccttgccaagtatatctgtgaaaa
 tcaagattcgatctccagtaaaactgaaggaatgctgtgaaaaacctctgt
 tggaaaaatcccactgcattgccgaagtggaaaaatgatgagatgcctgct
 gacttgccttcattagctgctgattttgttgaaagtaaggatgtttgcaa
 aaactatgctgaggcaaaggatgtcttctcctgggcatgtttttgtatgaat
 30 atgcaagaaggcatcctgattactctgtcgtgctgctgctgagacttgcc
 aagacatatgaaaccactctagagaagtgtgtgccgctgcagatcctca
 tgaatgctatgccaaagtgttcgatgaatttaaacctcttggtggaagagc
 ctcagaatttaatacaacaaaattgtgagctttttgagcagcttgagagag
 taaaaattccagaatgcgctattagttcgttacaccaagaaagtacccca
 35 agtgtcaactccaactcttgtagaggtctcaagaaacctagggaaaagtgg
 gcagcaaatgttgtaaacatcctgaagcaaaaagaatgcctgtgcagaa

gactatctatccgtggctcctgaaccagttatgtgtgttgcagagaaaaac
 gccagtaagtgcagagtcaccaaagctgcacagaatccttggatgaaca
 ggcgaccatgcttttcagctctggaagtcgatgaaacatacgttcccaa
 gagtttaatgctgaaacattcaccttccatgcagatatatgcacactttc
 5 tgagaaggagagacaaatcaagaaacaaactgcacttgttgagctcgtga
 aacacaagcccaaggcaacaaaagagcaactgaaagctgttatggatgat
 ttgcagctttttagagaagtgcagaggtgacgataaggagacctg
 ctttgccgaggagggtaaaaaacttgttgccgagtcaggccgccttag
 gcttaggatccccagctccagctccaggaagcgcgcgaacggagaccac
 10 tgtccgctcggggccggcggttgcctgcctgcacacggctccgcgcgtc
 gctggaagacctgggctgggcccattgggtgctgtcgccacgggaggtgc
 aagtgacctgtgcatcggcgctgcccagagccagttccggggcggaac
 atgcacgcgcagatcaagacgagcctgcaccgcctgaagcccgcacgggt
 gccagcgcctgtgctgctgcccgcagctacaatcccatggtgctcatc
 15 aaaagaccgacaccggggtgctgctccagacctatgatgacttgttagcc
 aaagactgccactgcatatga (SEQ ID NO:114).

As discussed above, in a specific embodiment, a homodimer is provided comprising two fusion proteins having the sequence of SEQ ID NO:115.

20 II.G.4 HSA-GS(PAPAP)₂GS- GDF15

The designation “HSA-GS(PAPAP)₂GS-GDF15” in the instant disclosure refers to a fusion protein comprising a GDF15 polypeptide linked to an HSA polypeptide via a linker comprising the sequence of SEQ ID NO:116 that connects the N-terminus of the GDF15 polypeptide to the C-terminus of the HSA polypeptide.

25 In certain embodiments, a homodimer is provided comprising two HSA-GS(PAPAP)₂GS-GDF15 fusion proteins, linked via an interchain disulfide bond between their respective GDF15 regions.

More particularly, in a specific embodiment, the homodimer comprises:

- (a) two HSA polypeptides (one each monomer) comprising the sequence of SEQ ID
 30 NO:110;
 (b) two GDF15 polypeptides (one each monomer) comprising the sequence of SEQ ID NO:12; and
 (c) two polypeptide linkers (one each monomer) comprising the sequence:

GSPAPAPPAPPGS (SEQ ID NO:116)

35 each linking the N-terminus of the GDF-15 polypeptide to the C-terminus of the HSA polypeptide via a peptide bond.

In a preferred embodiment, the fusion protein comprises the amino acid sequence (linker double underlined):

DAHKSEVAHRFKDLGEENFKALVLI~~AFAYLQQCPFEDHVKLVNEVTEFA~~
 KTCVADESAENCDKSLHTLFGDKLCTVATLRETYGEMADCCAKQEPERNE
 5 CFLQHKDDNP~~LPRLVRPEVDVMCTAFHDNEETFLKKYLYEIARRHPYFY~~
 APELLFFAKRYKAAFTECCQAADKAACLLPKLDEL~~RDEGKASSAKQRLKC~~
 ASLQKFGERAFKAWAVARLSQRFPKAEFAEVSKLVTDLT~~TKVHTECCHGDL~~
 LECADDRADLAKYICENQDSISSKLKECCEKPLLEKSHCIAEVENDEMPA
 DLPSLAADFVESKDVCKNYAEAKDVFLGMFLY~~EYARRHPDYSVVLRLLA~~
 10 KTYETTLEKCCAAADPHECYAKVFDEFKPLVEEPQNL~~IKQNC~~ELFEQLGE
 YKFQNALLVRYTKKVPQVSTPTLVEVSRNLGKVGSKCCKHPEAKRMPCAE
 DYLSVVLNQLCVLHEKTPVSDRVTKCCTESLVNRRPCFSALEVDETYVPK
 EFNAETFTFHADICTLSEKERQIKKQTALVELVKHKPKATKEQLKAVMDD
 FAAFVEKCKADDKETCFAEEGKKLVAASQAALGL~~GSPAPAPPAPAPGSA~~
 15 RNGDHCPLGPGRCCRLHTVRASLEDLGWADWVLSPREVQV~~TM~~CIGACPSQ
 FRAANMHAQIKTSLHRLKPDTVPAPCCVPASYNPMVLIQKTD~~TG~~VS~~LQ~~TY
 DDLLAKDCHCI (SEQ ID NO:118),

which is encoded by the nucleic acid sequence:

gatgcacacaagagtgaggttgctcatcgatttaaagatttgggagaaga
 20 aaatttcaaagccttggtggtgattgcctttgctcagtatcttcagcagt
 gtccatttgaagatcatgtaaaattagtgaaatgaagtaactgaatttgca
 aaaacatgtgttgctgatgagtcagctgaaaattgtgacaaatcacttca
 taccctttttggagacaaattatgcacagttgcaactcttcgtgaaacct
 atggtgaaatggctgactgctgtgcaaaacaagaacctgagagaaatgaa
 25 tgcttcttgcaacacaaagatgacaacccaaacctccccgattggtgag
 accagaggttgatgtgatgtgcaactgcttttcatgacaatgaagagacat
 ttttgaaaaaatacttatatgaaattgccagaagacatccttacttttat
 gccccggaactccttttctttgctaaaaggtataaagctgcttttacaga
 atgttgccaagctgctgataaagctgcctgcctgttgccaaagctcgatg
 30 aacttcgggatgaaggaaggcttcgtctgccaacagagactcaagtgt
 gccagtctccaaaaatttggagaaagagctttcaaagcatgggcagtagc
 tcgcctgagccagagatttcccaaagctgagtttgagaagtttccaagt
 tagtgacagatcttaccaaagtccacacggaatgctgcatggagatctg
 cttgaatgtgctgatgacagggcgaccttgccaagtatatctgtgaaaa
 35 tcaagattcgatctccagtaaactgaaggaatgctgtgaaaaacctctgt
 tggaaaaatcccactgcattgccgaagtggaaaaatgatgagatgcctgct

gacttgccttcattagctgctgattttgttgaaagtaaggatgtttgcaa
 aaactatgctgaggcaaaggatgtcttcctgggcatgtttttgtatgaat
 atgcaagaaggcatcctgattactctgtcgtgctgctgagacttgcc
 aagacatatgaaaccactctagagaagtgctgtgccgctgcagatcctca
 5 tgaatgctatgccaaagtgttcgatgaatttaaaccctcttggtgaagagc
 ctgagaatttaatacaacaaaattgtgagctttttgagcagcttgagag
 taaaaattccagaatgcgctattagttcgttacaccaagaaagtaccca
 agtgtcaactccaactcttgtagaggtctcaagaaacctagggaaaagtgg
 gcagcaaatgttgtaaacatcctgaagcaaaaagaatgccctgtgcagaa
 10 gactatctatccgtggctcctgaaccagttatgtgtgttgcatgagaaaac
 gccagtaagtgcagagtcaccaaagtctgcacagaatccttggtgaaca
 ggcgaccatgcttttcagctctggaagtcgatgaaacatacgttcccaa
 gagtttaatgctgaaacattcaccttccatgcagatatatgcacactttc
 tgagaaggagagacaaatcaagaaacaaactgcacttggtgagctcgtga
 15 aacacaagcccaaggcaacaaaagagcaactgaaagctgttatggatgat
 ttgcgagctttttgtagagaagtgctgcaaggctgacgataaggagacctg
 ctttgccgaggagggtaaaaaacttggtgcggccagtcaggccgccttag
 gcttaggatccccagctccagctccacccgcacctgcccctggaagcgcg
 cgcaacggagaccactgtccgctcggggcccggttgctgccgtctgca
 20 cacgggtccgcgctcgtggaagacctgggctgggcccattgggtgctgt
 cgccacgggaggtgcaagtgacctgtgcatcggcgctgcccagagccag
 ttccgggcggaacatgcacgcgcagatcaagacgagcctgcaccgcct
 gaagcccgacacggtgccagcgccctgctgcgtgcccgcagctacaatc
 ccatggtgctcattcaaaagaccgacaccggggtgtcgctccagacctat
 25 gatgacttggttagccaaagactgccactgcatatga (SEQ ID NO:117) .

As discussed above, in a specific embodiment, a homodimer is provided comprising two fusion proteins having the sequence of SEQ ID NO:118.

II.G.5 HSA-GSAAQAAQQGS-GDF15

30 The designation "HSA-GSAAQAAQQGS-GDF15 in the instant disclosure refers to a fusion protein comprising a GDF15 polypeptide linked to an HSA polypeptide via a linker comprising the sequence of SEQ ID NO:119 that connects the N-terminus of the GDF15 polypeptide to the C-terminus of the HSA polypeptide.

35 In certain embodiments, a homodimer is provided comprising two HSA-GSAAQAAQQGS-GDF15 fusion proteins, linked via an interchain disulfide bond between their respective GDF15 regions.

More particularly, in a specific embodiment, the homodimer comprises:

(a) two HSA polypeptides (one each monomer) comprising the sequence of SEQ ID NO:110;

(b) two GDF15 polypeptides (one each monomer) comprising the sequence of SEQ ID NO:12; and

(c) two polypeptide linkers (one each monomer) comprising the sequence:

GSAAQAAQQGS (SEQ ID NO:119)

each linking the N-terminus of the GDF15 polypeptide to the C-terminus of an HSA polypeptide via a peptide bond.

In a preferred embodiment, the fusion protein comprises the amino acid sequence (linker double underlined):

DAHKSEVAHREFKDLGEENFKALVLIIFAQYLQQCPFEDHVKLVNEVTEFA
KTCVADESAENCCKSLHTLFGDKLCTVATLRETYGEMADCCAKQEPERNE
CFLQHKDDNPPLPRLVRPEVDVMCTAFHDNEETFLKKYLYEIARRHPYFY
15 APELLFFAKRYKAAFTECCQAADKAACLLPKLDELRLDEGKASSAKQRLKC
ASLQKFGERAFAKAWAVARLSQRFPAEFAEVSKLVTDLTQVHTECCHGDL
LECADDRADLAKYICENQDSISSKLKECCEKPLEKSHCIAEVENDEMPA
DLPSLAADFVESKDVCKNYAEAKDVFLGMFLYEYARRHPDYSVVLRLA
KTYETTLEKCCAAADPHECYAKVFDEFKPLVEEPQNLIKQNCLEFEQLGE
20 YKFQNALLVRYTKKVPQVSTPTLVEVSRNLGKVGSKCKHPEAKRMPCAE
DYL SVVLNQLCVLHEKTPVSDRVTKCTESLVNRRPCFSALEVDETYVPK
EFNAETFTFHADICTLSEKERQIKKQTALVELVKHKPKATKEQLKAVMDD
FAAFVEKCKADDKETCFAEEGKKLVAAASQAALGLGSAAQAAQQGSARNG
DHCPLGPGRCCRLHTVRASLEDLGWADWVLSPREVQVTMCIGACPSQFRA
25 ANMHAQIKTSLHRLKPDTPAPCCVPASYNPMVLIQKTDGTGVSQTYDDL
LAKDCHCI (SEQ ID NO:121),

which is encoded by the nucleic acid sequence:

gatgcacacaagagtgaggttgctcatcgatttaaagatttgggagaaga
aaatttcaaagccttggtggttgattgcctttgctcagtatcttcagcagt
30 gtccatttgaagatcatgtaaaattagtgatgaagtaactgaatttgca
aaaacatgtgttgctgatgagtcagctgaaaattgtgacaaatcacttca
tacccttttttgagacaaattatgcacagttgcaactcttcgtgaaacct
atggtgaaatggctgactgctgtgcaaaacaagaacctgagagaaatgaa
tgcttcttgcaacacaaagatgacaacccaaacctccccgattggtgag
35 accagaggttgatgtgatgtgcactgcttttcatgacaatgaagagacat

ttttgaaaaataacttatatgaaattgccagaagacatccttacttttat
 gccccggaactccttttctttgctaaaaggtataaagctgcttttacaga
 atgttgccaagctgctgataaagctgcctgcctgttgccaaagctcgatg
 aacttcgggatgaagggaaggcttcgtctgccaaacagagactcaagtgt
 5 gccagtctccaaaaatttgagaaaagagctttcaaagcatgggcagtagc
 tcgcctgagccagagatttcccaaagctgagtttgagaagtttccaagt
 tagtgacagatcttaccaaagtcacacggaatgctgccatggagatctg
 cttgaatgtgctgatgacagggcgaccttgccaagtatatctgtgaaaa
 tcaagattcgatctccagtaaaactgaaggaatgctgtgaaaaacctctgt
 10 tggaaaaatcccactgcattgccgaagtggaaaaatgatgagatgcctgct
 gacttgccctcattagctgctgattttgttgaaagtaaggatgtttgcaa
 aaactatgctgaggcaaaggatgtcttcctgggcatgtttttgtatgaat
 atgcaagaaggcatcctgattactctgtcgtgctgctgctgagacttgcc
 aagacatatgaaaccactctagagaagtgtgtgccgctgcagatcctca
 15 tgaatgctatgccaaagtgttcgatgaatttaaaccctcttgtggaagagc
 ctgagaatttaatacaacaaaattgtgagctttttgagcagcttgagag
 tacaaattccagaatgcgctattagttcgttacaccaagaaagtaccca
 agtgtcaactccaactcttgttagaggtctcaagaaacctaggaaaagtgg
 gcagcaaatgttgtaaacatcctgaagcaaaaagaatgcctgtgcagaa
 20 gactatctatccgtggtcctgaaccagttatgtgtgttgcatgagaaaac
 gccagtaagtgcagagtcaccaaagtctgcacagaatccttggtgaaca
 ggcgaccatgcttttcagctctggaagtcgatgaaacatacgttcccaa
 gagtttaatgctgaaacattcaccttccatgcagatatatgcacactttc
 tgagaaggagagacaaatcaagaaacaaactgcacttggtgagctcgtga
 25 aacacaagcccaaggcaacaaaagagcaactgaaagctgttatggatgat
 ttgcgagctttttagagaagtgtgcaaggctgacgataaggagacctg
 ctttgccgaggagggtaaaaaacttggtgcggccagtcaggccgccttag
 gcttaggatccgccgctcaggctgcacagcaaggaagcgcgcgcaacgga
 gaccactgtccgctcgggcccgggcttgctgccgtctgcacacggtccg
 30 cgcgtcgttggaagacctgggctgggcccattgggtgctgtcgccacggg
 aggtgcaagtgacctgtgcatcggcgcgtgcccgagccagttccggggc
 gcaaacatgcacgcgcagatcaagacgagcctgcaccgcctgaagccga
 cacggtgccagcgcctgctgcgtgcccgccagctacaatcccatggtgc
 tcattcaaaagaccgacacggggtgtcgtccagacctatgatgacttg
 35 ttagccaaagactgccactgcatatga (SEQ ID NO:120).

As discussed above, in a specific embodiment, a homodimer is provided comprising two fusion proteins having the sequence of SEQ ID NO:121.

II.G.6 HSA-GS(AAQAAQQ)₂GS-GDF15

The designation “HSA-GS(AAQAAQQ)₂GS-GDF15” in the instant disclosure refers to a fusion protein comprising a GDF15 polypeptide linked to an HSA polypeptide via a linker comprising the sequence of SEQ ID NO:122 that connects the N-terminus of the GDF15 polypeptide to the C-terminus of the HSA polypeptide.

In certain embodiments, a homodimer is provided comprising two HSA-GS(AAQAAQQ)₂GS-GDF15 fusion proteins, linked via an interchain disulfide bond between their respective GDF15 regions.

More particularly, in a specific embodiment, the homodimer comprises:

(a) two HSA polypeptides (one each monomer) comprising the sequence of SEQ ID NO:110;

(b) two GDF15 polypeptides (one each monomer) comprising the sequence of SEQ ID NO:12; and

(c) two polypeptide linkers (one each monomer) comprising the sequence:

GSAAQAAQQAAQAAQQGS (SEQ ID NO:122)

each linking the N-terminus of the GDF-15 polypeptide to the C-terminus of the HSA polypeptide via a peptide bond.

In a preferred embodiment, the fusion protein comprises the amino acid sequence (linker double underlined):

DAHKSEVAHRFKDLGEENFKALVLI AFAQYLQQCPFEDHVKLVNEVTEFA
KTCVADESAENCDKSLHTLFGDKLCTVATLRETYGEMADCCAKQEPERNE
CFLQHKDDNPNLPRLVRPEVDVMCTAFHDNEETFLKKYLYEIARRHPYFY
APELLFFAKRYKAAFTECCQAADKAAACLLPKLDEL RDEGKASSAKQRLKC
25 ASLQKFGERAFKAWAVARLSQRFPKAEFAEVSKLVTDLT KVHTECCHGDL
LECADDRADLAKYICENQDSISSKLKECCEKPLEKSHCIAEVENDEMPA
DLPSLAADFVESKDVCKNYAEAKDVFLGMFLYEYARRHPDYSVVL LRLA
KTYETTLEKCCAAADPHECYAKVFDEFKPLVEEPQNLIKQNC ELFQQLGE
YKFQNALLVRYTKKVPQVSTPTLVEVSRNLGKVGSKCKHPEAKRMPCAE
30 DYLSVVLNQLCVLHEKTPVSDRVTKCCTESLVNRRPCFSALEVD ETYVPK
EFNAETFTFHADICTLSEKERQIKKQTALVELVKHKPKATKEQLKAVMDD
FAAFVEKCKADDKETCFAEEGKKLVAAASQAALGLGSAAQAAQQAQAAQ
QGSARNGDHCPLGPGRCCRLHTVRASLEDLGWADWVLSPREVQVTMCIGA
QGSARNGDHCPLGPGRCCRLHTVRASLEDLGWADWVLSPREVQVTMCIGA
CPSQFRAANMHAQIKTSLHRLKPDTPAPCCVPASYNPMVLIQKTD TGVS
35 LQTYDDLAKDCHCI (SEQ ID NO:124),

which is encoded by the nucleic acid sequence:

```

gatgcacacaagagtgaggttgctcatcgatttaaagatttgggagaaga
aaatttcaaaagccttggtggttgattgcctttgctcagtatcttcagcagt
gtccatttgaagatcatgtaaaattagtgaatgaagtaactgaatttgca
5  aaaacatgtgttgctgatgagtcagctgaaaattgtgacaaatcacttca
tacccttttttgagacaaattatgcacagttgcaactcttcgtgaaacct
atggtgaaatggctgactgctgtgcaaaacaagaacctgagagaaatgaa
tgcttcttgcaacacaaagatgacaacccaaacctccccgattggtgag
accagaggttgatgtgatgtgcactgcttttcatgacaatgaagagacat
10  ttttgaaaaaatacttatatgaaattgccagaagacatccttacttttat
gccccggaactccttttcttgctaaaaggtataaagctgcttttacaga
atggttgccaagctgctgataaagctgcctgcctgttgccaaagctcgatg
aacttcgggatgaagggaaggcttcgtctgccaacagagactcaagtgt
gccagtctccaaaaatttggagaaagagctttcaaagcatgggcagtagc
15  tcgcctgagccagagatttcccaaagctgagtttgagaagtttccaagt
tagtgacagatcttaccaaagtcacacggaatgctgccatggagatctg
cttgaatgtgctgatgacagggcggaaccttgccaagtatatctgtgaaaa
tcaagattcgatctccagtaaaactgaaggaatgctgtgaaaaacctctgt
tggaaaaatcccactgcattgccgaagtggaaaaatgatgagatgcctgct
20  gacttgcccttcattagctgctgattttgttgaaagtaaggatgtttgcaa
aaactatgctgaggcaaaggatgtcttcctgggcatgtttttgtatgaat
atgcaagaaggcatcctgattactctgtcgtgctgctgctgagacttgcc
aagacatatgaaaccactctagagaagtgctgtgccgctgcagatcctca
tgaatgctatgccaaagtgttcgatgaatttaaacctcttgtggaagagc
25  ctcagaatttaatacaacaaaattgtgagctttttgagcagcttgagagag
tacaaattccagaatgcgctattagttcgttacaccaagaaagtaccca
agtggtcaactccaactcttgtagaggtctcaagaaacctaggaaaagtgg
gcagcaaatgttgtaaacatcctgaagcaaaaagaatgcctgtgcagaa
gactatctatccgtggtcctgaaccagttatgtgtgttgcatgagaaaac
30  gccagtaagtgcagagtcaccaaattgctgcacagaatccttgggtgaaca
ggcgaccatgcttttcagctctggaagtcgatgaaacatacgttcccaaa
gagtttaaatgctgaaacattcaccttccatgcagatatatgcacactttc
tgagaaggagagacaaatcaagaaacaaactgcacttggtgagctcgtga
aacacaagcccaaggcaacaaaagagcaactgaaagctgttatggatgat
35  ttgcgagctttttagagaagtgctgcaaggctgacgataaggagacctg
ctttgccgaggagggtaaaaaacttggtgcggccagtcaggccgccttag
gcttaggatccgccgctcaggctgcacagcaagcagcccaagcagctcag

```

caggggaagcgcgcgcaacggagaccactgtccgctcgggccccgggcgttg
 ctgccgtctgcacacgggtccgcgcgtcgctggaagacctgggctgggccc
 attgggtgctgtcgccacgggaggtgcaagtgacctgtgcatcggcgcg
 tgcccagagccagttccgggcggaacatgcacgcgcagatcaagacgag
 5 cctgcaccgcctgaagccccgacacgggtgccagcgccctgctgcgtgccc
 ccagctacaatcccatggtgctcattcaaaagaccgacaccggggtgctg
 ctccagacctatgatgacttgtagccaaagactgccactgcatatga (SEQ ID
 NO:123) .

As discussed above, in a specific embodiment, a homodimer is provided comprising
 10 two fusion proteins having the sequence of SEQ ID NO:124.

II.G.7 HSA-GGNAEAAAKEAAAKEAAKAGG-GDF15

The designation “HSA-GGNAEAAAKEAAAKEAAKAGG-GDF15” in the instant
 disclosure refers to a fusion protein comprising a GDF15 polypeptide linked to an HSA
 15 polypeptide via a linker comprising the sequence of SEQ ID NO:125 that connects the
 N-terminus of the GDF15 polypeptide to the C-terminus of the HSA polypeptide.

In certain embodiments, a homodimer is provided comprising two HSA-
 GGNAEAAAKEAAAKEAAKAGG-GDF15 fusion proteins, linked via an interchain
 disulfide bond between their respective GDF15 regions.

More particularly, in a specific embodiment, the homodimer comprises:
 (a) two HSA polypeptides (one each monomer) comprising the sequence of SEQ ID
 NO:110;
 (b) two GDF15 polypeptides (one each monomer) comprising the sequence of SEQ
 ID NO:12; and
 25 (c) two polypeptide linkers (one each monomer) comprising the sequence:

GGNAEAAAKEAAAKEAAKAGG (SEQ ID NO:125)

each linking the N-terminus of the GDF-15 polypeptide to the C-terminus of the HSA
 polypeptide via a peptide bond.

In a preferred embodiment, the fusion protein comprises the amino acid sequence
 30 (linker double underlined):

DAHKSEVAHREFKDLGEENFKALVLIAFAQYLQQCPFEDHVKLNVNEVTEFA
 KTCVADESAENCDKSLHTLFGDKLCTVATLRETYGEMADCCAKQEPERNE
 CFLQHKDDNPNLRLVRPEVDVMCTAFHDNEETFLKKYLYEIARRHPYFY
 APELLFFAKRYKAAFTECCQAADKAACLLPKLDELRLDEGKASSAKQRLKC
 35 ASLQKFGERAFAKAWAVARLSQRFPAEFAEVSKLVTDLTKVHTECCHGDL

LECADDRADLAKYICENQDSISSKLKECCEKPLLEKSHCIAEVENDEMPA
 DLPSLAADFVESKDVCKNYAEAKDVFLGMFLYEYARRHPDYSVVLRLA
 KTYETTLEKCCAAADPHECYAKVFDEFKPLVEEPQNLIKQNCSELFELGE
 YKFQNALLVRYTKKVPQVSTPTLVEVSRNLGKVGSKCKHPEAKRMPCAE
 5 DYLSVVLNQLCVLHEKTPVSDRVTKCTESLVNRRPCFSALEVDETYVPK
 EFNAETFTFHADICTLSEKERQIKKQTALVELVKHKPKATKEQLKAVMDD
 FAAFVEKCKKADDKETCFAEEGKKLVAASQAALGLGGNAEAAAKEAAAKE
AAAKEAAAKAGGARNGDHCPLGPGRCRLHTVRASLEDLGWADWVLSPRE
 VQVTMCIGACPSQFRAANMHAQIKTSLHRLKPDTVPAPCCVPASYNPMVL
 10 IQKTDGTGVSLLQTYDDLAKDCHCI (SEQ ID NO:127),

which is encoded by the nucleic acid sequence:

gatgcacacaagagtgaaggtgctcatcgatttaaagatttgggagaaga
 aaatttcaaagccttggtgttgattgccttgcctcagtatcttcagcagt
 gtccatttgaagatcatgtaaaattagtgaatgaagtaactgaatttgca
 15 aaaacatgtgttgctgatgagtcagctgaaaattgtgacaaatcacttca
 taccctttttggagacaaattatgcacagttgcaactcttcgtgaaacct
 atggtgaaatggctgactgctgtgcaaaacaagaacctgagagaaatgaa
 tgcttcttgcaacacaaagatgacaacccaaacctccccgattggtgag
 accagaggttgatgtgatgtgactgcttttcatgacaatgaagagacat
 20 ttttgaaaaaataacttatatgaaattgccagaagacatccttacttttat
 gccccggaactccttttcttgctaaaaggtataaagctgcttttacaga
 atgttgccaagctgctgataaagctgcctgcctgttgccaaagctcgatg
 aacttcgggatgaagggaaggcttcgtctgccaacagagactcaagtgt
 gccagtcctccaaaatttggagaaagagctttcaaagcatgggcagtagc
 25 tcgcctgagccagagatttcccaaagctgagtttgagaagtttccaagt
 tagtgacagatcttaccaaagtcacacggaatgctgccatggagatctg
 cttgaatgtgctgatgacagggcggaaccttgccaagtatatctgtgaaaa
 tcaagattcgatctccagtaaaactgaaggaatgctgtgaaaaacctctgt
 tggaaaaatcccactgcattgccgaagtggaaaaatgatgagatgcctgct
 30 gacttgccttcattagctgctgattttgttgaaagtaaggatgtttgcaa
 aaactatgctgaggcaaaggatgtcttcctgggcatgtttttgtatgaat
 atgcaagaaggcatcctgattactctgtcgtgctgctgctgagacttgcc
 aagacatatgaaaccactctagagaagtgctgtgccgctgcagatcctca
 tgaatgctatgccaaagtgttcgatgaatttaaacctcttggtggaagagc
 35 ctgagaatttaatacaacaaaattgtgagctttttgagcagcttgagagag
 taaaaattccagaatgcgctattagttcggttacaccaagaaagtacccca

agtgtcaactccaactcttgttagaggtctcaagaaacctaggaaaagtgg
 gcagcaaatgttgtaaacatcctgaagcaaaaagaatgcctgtgcagaa
 gactatctatccgtggctcctgaaccagttatgtgtgttgcatgagaaaac
 gccagtaagtgcagagtcaccaaagtctgcacagaatccttggtgaaca
 5 ggcgaccatgcttttcagctctggaagtcgatgaaacatacgttcccaa
 gagtttaatgctgaaacattcaccttccatgcagatatatgcacactttc
 tgagaaggagagacaaatcaagaaacaaactgcacttggtgagctcgtga
 aacacaagcccaaggcaacaaaagagcaactgaaagctgttatggatgat
 ttcgcagctttttagagagaagtgtgcaaggctgacgataaggagacctg
 10 ctttgccgaggagggtaaaaaacttggtgcgccagtcaggccgccttag
 gcttaggaggcaacgccgaggctgccgctaaggaagccgctgccaaggag
 gccgcagcaaaaagaggctgcagctaaggccggaggagcgcgcaacggaga
 ccactgtccgctcggggccggcggttgctgccgtctgcacacggtccgcg
 cgctcgctggaagacctgggctgggcccagattgggtgctgtcgccacgggag
 15 gtgcaagtgaccatgtgcatcggcgcgtgcccgagccagttccgggcggc
 aaacatgcacgcgcagatcaagacgagcctgcaccgctgaagcccgcaca
 cgggtgccagcgccctgctgcgtgcccgccagctacaatcccatgggtgctc
 attcaaaagaccgacaccggggtgctcgctccagacctatgatgacttggt
 agccaaagactgccactgcatatga (SEQ ID NO:126) .

20 As discussed above, in a specific embodiment, a homodimer is provided comprising
 two fusion proteins having the sequence of SEQ ID NO:127.

II.G.8 HSA-(G₄S)₆-GDF15

The designation “HSA-(G₄S)₆-GDF15” in the instant disclosure refers to a fusion
 25 protein comprising a GDF15 polypeptide linked to an HSA polypeptide via a linker
 comprising the sequence of SEQ ID NO:128 that connects the N-terminus of the GDF15
 polypeptide to the C-terminus of the HSA polypeptide.

In certain embodiments, a homodimer is provided comprising two HSA-(G₄S)₆-
 GDF15 fusion proteins, linked via an interchain disulfide bond between their respective
 30 GDF15 regions.

More particularly, in a specific embodiment, the homodimer comprises:

- (a) two HSA polypeptides (one each monomer) comprising the sequence of SEQ ID
 NO:110;
- (b) two GDF15 polypeptides (one each monomer) comprising the sequence of SEQ
 35 ID NO:12; and
- (c) two polypeptide linkers (one each monomer) comprising the sequence:

tcgcctgagccagagagatttcccaaagctgagtttgagaagtttccaagt
 tagtgacagatcttaccaaagtcacacggaatgctgcatggagatctg
 cttgaatgtgctgatgacagggcggaccttgccaagtatatctgtgaaa
 tcaagattcgatctccagtaaaactgaaggaatgctgtgaaaaacctctgt
 5 tggaaaaatcccactgcattgccgaagtggaaaatgatgagatgcctgct
 gacttgcttcattagctgctgattttgttgaaagtaaggatgtttgcaa
 aaactatgctgaggcaaaggatgtcttcctgggcatgtttttgtatgaat
 atgcaagaaggcatcctgattactctgtcgtgctgctgctgagacttgcc
 aagacatatgaaaccactctagagaagtgctgtgccgctgcagatcctca
 10 tgaatgctatgccaaagtgttcgatgaatttaaacctcttggtggaagagc
 ctgagaatttaatacaaaaaattgtgagctttttgagcagcttgagag
 taaaaattccagaatgcgctattagttcgttacaccaagaaagtaccca
 agtgtcaactccaactcttgtagaggtctcaagaaacctaggaaaagtgg
 gcagcaaatgttgtaaacatcctgaagcaaaaagaatgccctgtgcagaa
 15 gactatctatccgtggtcctgaaccagttatgtgtgttgcatgagaaaac
 gccagtaagtgcagagtcaccaaagtctgcacagaatccttggtgaaca
 ggcgaccatgcttttcagctctggaagtcgatgaaacatacgttcccaa
 gagtttaaatgctgaaacattcaccttccatgcagatatatgcacactttc
 tgagaaggagagacaaatcaagaaacaaactgcacttgttgagctcgtga
 20 aacacaagcccaaggcaacaaaagagcaactgaaagctgttatggatgat
 ttgcgagctttttgtagagaagtgctgcaaggctgacgataaggagacctg
 ctttgccgaggagggtaaaaaacttgttgcgccagtcaggccgccttag
 gcttaggaggtggtggtctctggaggcggtggaagcggaggcggtggatcc
 ggaggcggtggaagcggaggtggtggatctggaggcggtggaagcgcgcg
 25 caacggagaccactgtccgctcgggcccggcggttgctgccgtctgcaca
 cgggtccgcgcgtcgtggaagacctgggctgggccgattgggtgctgtcg
 ccacgggaggtgcaagtgacctgtgcatcggcgcgtgcccagagccagtt
 ccgggcggaacatgcacgcgcagatcaagacgagcctgcaccgcctga
 agcccagacacggtgccagcgccctgctgcgtgcccgcagctacaatccc
 30 atggtgctcattcaaaagaccgacacccgggtgtcgtccagacctatga
 tgacttggttagccaaagactgccactgcatatga (SEQ ID NO:129).

As discussed above, in a specific embodiment, a homodimer is provided comprising two fusion proteins having the sequence of SEQ ID NO:130.

35 II.G.9 HSA-GS(AAQAAQQ)₂GS-GDF15(N3D)

The designation “HSA-GS(AAQAAQQ)₂GS-GDF15(N3D)” in the instant disclosure refers to a fusion protein comprising a GDF15(N3D) polypeptide linked to an HSA

polypeptide via a linker comprising the sequence of SEQ ID NO:122 that connects the N-terminus of the GDF15(N3D) polypeptide to the C-terminus of the HSA polypeptide.

In certain embodiments, a homodimer is provided comprising two HSA-GS(AAQAAQQ)₂GS-GDF15(N3D) fusion proteins, linked via an interchain disulfide bond
5 between their respective GDF15 regions.

More particularly, in a specific embodiment, the homodimer comprises:

(a) two HSA polypeptides (one each monomer) comprising the sequence of SEQ ID NO:110;

(b) two GDF15(N3D) polypeptides (one each monomer) comprising the sequence of
10 SEQ ID NO:52; and

(c) two polypeptide linkers (one each monomer) comprising the sequence of SEQ ID NO:122 each linking the N-terminus of a GDF15(N3D) polypeptide to the C-terminus of an HSA polypeptide via a peptide bond.

In a preferred embodiment, the fusion protein comprises the amino acid sequence
15 (linker double underlined):

DAHKSEVAHRFKDLGEENFKALVLIIFAQYLQQCPFEDHVKLNVNEVTEFA
KTCVADESAENCDSLHTLFGDKLCTVATLRETYGEMADCCAKQEPERNE
CFLQHKDDNPPLRLVRPEVDVMCTAFHDNEETFLKKYLYEIARRHPYFY
APELLFFAKRYKAAFTECCQAADKAACLLPKLDELRLDEGKASSAKQRLKC
20 ASLQKFGERAFAKAWAVARLSQRFPKAEFAEVSKLVTDLTKVHTECCHGDL
LECADDRADLAKYICENQDSISSKLKECCEKPLEKSHCIAEVENDEMPA
DLPSLAADFVESKDVCKNYAEAKDVFLGMFLYEYARRHPDYSVVLRLRLA
KTYETTLEKCCAAADPHECYAKVFDEFKPLVEEPQNLIKQNCSELFQQLGE
YKFQNALLVRYTKKVPQVSTPTLVEVSRNLGKVGSKCKHPEAKRMPCAE
25 DYLSVVLNQLCVLHEKTPVSDRVTKCTESLVNRRPCFSALEVDETYVPK
EFNAETFTFHADICTLSEKERQIKKQTALVELVKHKPKATKEQLKAVMDD
FAAFVEKCKKADDKETCFAEEGKKLVAASQAALGLGSAAQAAQAAQAAQ
QGSARDGDHCPGLGPRCCRLHTVRASLEDLGWADWVLSPREVQVTMCIGA
CPSQFRAANMHAQIKTSLHRLKPDTVPAPCCVPASYNPMVLIQKTDGTGVS
30 LQTYDDLAKDCHCI (SEQ ID NO:242),

which is encoded by the nucleic acid sequence:

gatgcacacaagagtggaggttgctcatcgatttaaagatttg
gagaagaaaatttcaaagccttggtggtgattgcctttgctcagtatctt
cagcagtggtccatttgaagatcatgtaaaattagtgaatgaagtaactga
35 atttgcaaaaacatgtgttgctgatgagtcagctgaaaattgtgacaaat
cacttcataaccctttttggagacaaattatgcacagttgcaactcttcgt

gaaacctatggtgaaatggctgactgctgtgcaaaacaagaacctgagag
aatgaatgcttcttgcaacacaaagatgacaacccaaacctccccgat
tggtgagaccagaggttgatgtgatgtgcaactgcttttcatgacaatgaa
gagacatTTTTTgaaaaatacttatatgaaattgccagaagacatcetta
5 cttttatgccccggaactccttttctttgctaaaagggtataaagctgctt
ttacagaatgttgccaagctgctgataaagctgcctgcctgttgccaaag
ctcgatgaacttcgggatgaagggaaggcttcgtctgccaaacagagact
caagtgtgccagtctccaaaaatttgagagaaagagctttcaaagcatggg
cagtagctcgctgagccagagatttcccaaagctgagtttgcagaagtt
10 tccaagttagtgacagatcttaccaaagtcacacggaatgctgccatgg
agatctgcttgaatgtgctgatgacagggcggaccttgccaagtatatct
gtgaaaatcaagattcgatctccagtaaaactgaaggaatgctgtgaaaa
cctctgttgaaaaatcccactgcattgccgaagtggaaaatgatgagat
gcctgctgacttgccctcattagctgctgattttgttgaaagtaaggatg
15 tttgcaaaaactatgctgaggcaaaggatgtcttcctgggcatgtttttg
tatgaatatgcaagaaggcatcctgattactctgtcgtgctgctgctgag
acttgccaagacatatgaaaccactctagagaagtgtgtgccgctgcag
atcctcatgaatgctatgccaaagtgttcgatgaatttaaacctcttgtg
gaagagcctcagaatttaatacaaaaaattgtgagctttttgagcagct
20 tggagagtacaaattccagaatgcgctattagttcgttacaccaagaaaag
taccccaagtgtcaactccaactcttgtagaggtctcaagaaacctagga
aaagtgggcagcaaatgttgtaaacatcctgaagcaaaaagaatgccttg
tgcagaagactatctatccgtggctcctgaaccagttatgtgtgttgcatg
agaaaacgccagtaagtgcagagtcaccaaagtgtgcacagaatccttg
25 gtgaacaggcgaccatgcttttcagctctggaagtcgatgaaacatacgt
tcccaaagagtttaatgctgaaacattcaccttccatgcagatatatgca
cactttctgagaaggagagacaaatcaagaaacaaactgcacttggtgag
ctcgtgaaacacaagcccaaggcaacaaaagagcaactgaaagctgttat
ggatgatttcgcagctttttgtagagaagtgtgcaaggctgacgataagg
30 agacctgctttgccgaggagggtaaaaaacttggtgcggccagtcaggcc
gccttaggcttaggatccgcgctcaggctgcacagcaagcagcccaagc
agctcagcagggaagcgcgcgacggagaccactgtccgctcgggcccg
ggcgttgctgccgtctgcacacgggtccgcgctcgtggaagacctgggc
tgggcccgattgggtgctgtcgccacgggaggtgcaagtgaccatgtgcat
35 cggcgctgcccagccagttccgggcccgaacatgcacgcgcagatca
agacgagcctgcaccgcctgaagcccagacggtgccagcgcctgctgc
gtgcccgcagctacaatcccatggtgctcattcaaaagaccgacaccgg

gggtgtcgctccagacctatgatgacttggttagccaaagactgccactgca
ta (SEQ ID NO:241).

As discussed above, in a specific embodiment, a homodimer is provided comprising two fusion proteins having the sequence of SEQ ID NO:242.

5

II.H. Constructs with Mutations to Address Affinity for FcγR Binding

It was found that certain charged pair (delHinge) multimers exhibited Fcγ receptor (FcγR) binding, particularly to FcγRI and FcγRIII. See, e.g., Example 7. In some cases, the FcγR binding affinity was comparable to that of observed in multimers comprising a hinge region. This was unexpected, because the Fcγ receptor interacts with the hinge region and these multimers were delHinge, as described above, lacking all or part of the hinge region. Mutational analyses of the Fc residues involved in binding to FcγR suggest that the main interaction site is localized in the hinge region and CH2 domain (Tamm A, 1997, *Int. Rev. Immunol.* 16:57–85). See also, Radaev S, et al., *J. Biol. Chem.* 276:16469–16477; Sondermann, P, et al., 2000, *Nature* 406:267–273.

Antibody-dependent cellular cytotoxicity (ADCC), an immune response mediated primarily by natural killer (NK) cells in humans, depends on the interaction of FcγRs, especially FcγRIIIA in humans, with the Fc domain of an antibody or Fc-containing protein. In ADCC, Fc binding to FcγRIII on the surface of an NK cell activates the NK cell, which releases perforins and granzymes.

Accordingly, constructs are provided having additional modifications to modulate the interaction of the construct with the FcγR. In one series of embodiments, an asparagine to glycine mutation (N297G) is introduced to a native Fc or Fc variant, including the various Fc domains described above. The N297G mutation removes a conserved N-glycosylation site at in the CH2 domain.

In another series of embodiments, additional N-terminal amino acid residues are deleted from an Fc domain from which all or a portion of the hinge region has been removed. For example, the amino acid sequence:

GGPSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVDGVEVH
NAKTKPREEQYNSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPAPIEKT
ISKAKGQPREPQVYTLPPSREEMTKNQVSLTCLVKGFYPSDIAVEWESNG
QPENNYKTTTPVLDSDGSFFLYSKLTVDKSRWQQGNVFSVSMHEALHNH
YTQKSLSLSPGK (SEQ ID NO:303)

is provided by deletion of amino acid residues N-terminal to G236 of the Fc domain of wild-type IgG1. In some embodiments, the C-terminal lysine (K447) optionally may be deleted from this Fc variant. The amino acid sequence:

GPSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVDGVEVHN
 5 AKTKPREEQYNSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPAPIEKT
 SKAKGQPREPQVYTLPPSREEMTKNQVSLTCLVKGFYPSDIAVEWESNGQ
 PENNYKTTTPVLDSGDSFFLYSKLTVDKSRWQQGNVFSVMSHEALHNHY
 TQKSLSLSPGK (SEQ ID NO: 304)

is provided by deletion of amino acid residues N-terminal to G237 from the Fc domain of wild-type IgG1. In some embodiments, the C-terminal lysine (K447) optionally may be deleted from this Fc variant. The amino acid sequence:

PSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVDGVEVHNA
 KTKPREEQYNSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPAPIEKTIS
 KAKGQPREPQVYTLPPSREEMTKNQVSLTCLVKGFYPSDIAVEWESNGQP
 15 ENNYKTTTPVLDSGDSFFLYSKLTVDKSRWQQGNVFSVMSHEALHNHYT
 QKSLSLSPGK (SEQ ID NO: 305)

is provided by deletion of amino acid residues N-terminal to P238 of the Fc domain of wild-type IgG1. In some embodiments, the C-terminal lysine (K447) optionally may be deleted from this Fc variant.

20 In one embodiment, a DhCpmFc(-) domain into which an asparagine to glycine mutation (N297G) has been introduced is provided (referred to herein as a “DhCpmFc(-)(N297G)” domain). In another embodiment, a DhCpmFc(+) domain into which an asparagine to glycine mutation (N297G) has been introduced is provided (referred to herein as a “DhCpmFc(+)(N297G)” domain).

25 In another embodiment, a DhCpmFc(-)(Y349C) domain into which an asparagine to glycine mutation (N297G) has been introduced is provided (referred to herein as a “DhCpmFc(-)(N297G)(Y349C)” domain). In another embodiment, a DhCpmFc(+)(S354C) domain into which an asparagine to glycine mutation (N297G) has been introduced is provided (referred to herein as a “DhCpmFc(+)(N297G)(S354C)” domain).

30 In another embodiment, a DhCpmFc(+)(Y349C) domain into which an asparagine to glycine mutation (N297G) has been introduced is provided (referred to herein as a “DhCpmFc(+)(N297G)(Y349C)” domain). In another embodiment, a DhCpmFc(-)(S354C) domain into which an asparagine to glycine mutation (N297G) has been introduced is provided (referred to herein as “DhCpmFc(-)(N297G)(S354C)” domain).

In another embodiment, a DhCpmFc(+)(L351C) domain into which an asparagine to glycine mutation (N297G) has been introduced is provided (referred to herein as a “DhCpmFc(+)(N297G)(L351C)” domain). In another embodiment, a DhCpmFc(-)(L351C) domain into which an asparagine to glycine mutation (N297G) has been introduced is provided (referred to herein as a “DhCpmFc(-)(N297G)(L351C)” domain).

In another embodiment, a DhCpmFc(-) domain into which an alanine to cysteine mutation (A287C) has been introduced is provided (referred to herein as a “DhCpmFc(-)(A287C)” domain). In another embodiment, a DhCpmFc(+) domain into which an alanine to cysteine mutation (A287C) has been introduced is provided (referred to herein as a “DhCpmFc(+)(A287C)” domain).

In another embodiment, a DhCpmFc(-) domain into which a leucine to cysteine mutation (L306C) has been introduced is provided (referred to herein as a “DhCpmFc(-)(L306C)” domain). In another embodiment, a DhCpmFc(+) domain into which a leucine to cysteine mutation (L306C) has been introduced is provided (referred to herein as a “DhCpmFc(+)(L306C)” domain).

In another embodiment, a DhCpmFc(-)(A287C) domain into which a tyrosine to cysteine mutation (Y349C) has been introduced is provided (referred to herein as a “DhCpmFc(-)(A287C)(Y349C)” domain). In another embodiment, a DhCpmFc(+)(A287C) domain into which a tyrosine to cysteine mutation (Y349C) has been introduced is provided (referred to herein as a “DhCpmFc(+)(A287C)(Y349C)” domain).

In another embodiment, a DhCpmFc(-)(A287C) domain into which a serine to cysteine mutation (S354C) has been introduced is provided (referred to herein as a “DhCpmFc(-)(A287C)(S354C)” domain). In another embodiment, a DhCpmFc(+)(A287C) domain into which a serine to cysteine mutation (S354C) has been introduced is provided (referred to herein as a “DhCpmFc(+)(A287C)(S354C)” domain).

In another embodiment, a DhCpmFc(-)(L306C) domain into which a tyrosine to cysteine mutation (Y349C) has been introduced is provided (referred to herein as a “DhCpmFc(-)(L306C)(Y349C)” domain). In another embodiment, a DhCpmFc(+)(L306C) domain into which a tyrosine to cysteine mutation (Y349C) has been introduced is provided (referred to herein as a “DhCpmFc(+)(L306C)(Y349C)” domain).

In another embodiment, a DhCpmFc(-)(L306C) domain into which a serine to cysteine mutation (S354C) has been introduced is provided (referred to herein as a “DhCpmFc(-)(L306C)(S354C)” domain). In another embodiment, a DhCpmFc(+)(L306C) domain into which a serine to cysteine mutation (S354C) has been introduced is provided (referred to herein as a “DhCpmFc(+)(A287C)(L306C)” domain).

In another embodiment, a DhCpmFc(-) domain from which the N-terminal 7 amino acids have been removed is provided (referred to herein as a “Dh2CpmFc(-)” domain). In

another embodiment, a DhCpmFc(+) domain from which the N-terminal 7 amino acids have been removed is provided (referred to herein as a “Dh2CpmFc(+)” domain).

In another embodiment, a DhCpmFc(-)(Y349C) domain from which the N-terminal 7 amino acids have been removed is provided (referred to herein as a “Dh2CpmFc(-)(Y349C)” domain). In another embodiment, a DhCpmFc(+)(S354C) domain from which the N-terminal 7 amino acids have been removed is provided (referred to herein as a “Dh2CpmFc(+)(S354C)” domain).

In another embodiment, a DhCpmFc(+)(Y349C) domain from which the N-terminal 7 amino acids have been removed is provided (referred to herein as a “Dh2CpmFc(+)(Y349C)” domain). In another embodiment, a DhCpmFc(-)(S354C) domain from which the N-terminal 7 amino acids have been removed is provided (referred to herein as a “Dh2CpmFc(-)(S354C)” domain).

In another embodiment, a CpmFc(+) domain into which an asparagine to glycine mutation (N297G) has been introduced is provided (referred to herein as a “CpmFc(+)(N297G)” domain). In another embodiment, a CpmFc(-) domain into which an asparagine to glycine mutation (N297G) has been introduced is provided (referred to herein as a “CpmFc(-)(N297G)” domain).

In another embodiment, a Dh2CpmFc(+) domain into which an asparagine to glycine mutation (N297G) has been introduced is provided (referred to herein as a “Dh2CpmFc(+)(N297G)” domain). In another embodiment, a Dh2CpmFc(-) domain into which an asparagine to glycine mutation (N297G) has been introduced is provided (referred to herein as a “Dh2CpmFc(-)(N297G)” domain).

In another embodiment, a Dh2CpmFc(-)(N297G) domain into which an alanine to cysteine mutation (A287C) has been introduced is provided (referred to herein as a “Dh2CpmFc(-)(N297G)(A287C)” domain). In another embodiment, Dh2CpmFc(+)(N297G) domain into which a leucine to cysteine mutation (L306C) has been introduced is provided (referred to herein as a “Dh2CpmFc(+)(N297G)(L306C)” domain).

In another embodiment, a Dh2CpmFc(-)(N297G)(A287C) domain into which a tyrosine to cysteine mutation (Y349C) has been introduced is provided (referred to herein as a “Dh2CpmFc(-)(N297G)(A287C)(Y349C)” domain). In another embodiment, a Dh2CpmFc(+)(N297G)(L306C) domain into which a serine to cysteine mutation (S354C) has been introduced is provided (referred to herein as a “Dh2CpmFc(+)(N297G)(L306C)(S354C)” domain).

In another embodiment, a Dh2CpmFc(-) domain onto which two glycines have been added to the N-terminus is provided (referred to herein as a “GG-Dh2CpmFc(-)(N297G)” domain). In another embodiment, a Dh2CpmFc(+) domain onto which two glycines have

been added to the N-terminus is provided (referred to herein as a “GG-Dh2CpmFc(+)(N297G)” domain).

In another embodiment, a GG-Dh2CpmFc(-) domain into which a tyrosine to cysteine mutation (Y349C) have been introduced is provided (referred to herein as a “GG-Dh2CpmFc(-)(Y349C)” domain). In another embodiment, a GG-Dh2CpmFc(+) domain into which a serine to cysteine mutation (S354C) has been introduced is provided (referred to herein as a “GG-Dh2CpmFc(+)(S354C)” domain).

In another embodiment, a GG-Dh2CpmFc(+) domain into which a tyrosine to cysteine mutation (Y349C) has been introduced is provided (referred to herein as a “GG-Dh2CpmFc(+)(Y349C)” domain). In another embodiment, a GG-Dh2CpmFc(-) domain into which a serine to cysteine mutation (S354C) has been introduced is provided (referred to herein as a “GG-Dh2CpmFc(-)(S354C)” domain).

In another embodiment, a Dh2CpmFc(-) domain onto which a glycine has been added to the N-terminus is provided (referred to herein as a “Dh3CpmFc(-)(N297G)” domain). In another embodiment, a Dh2CpmFc(+) domain onto which a glycine has been added to the N-terminus is provided (referred to herein as a “Dh3CpmFc(+)(N297G)” domain).

In another embodiment, a Dh3CpmFc(-) domain into which a tyrosine to cysteine mutation (Y349C) has been introduced is provided (referred to herein as a “Dh3CpmFc(-)(Y349C)” domain). In another embodiment, a Dh3CpmFc(+) domain into which a serine to cysteine mutation (S354C) has been introduced is provided (referred to herein as a “Dh3CpmFc(+)(S354C)” domain).

In another embodiment, an asparagine to glycine mutation (N297G) is introduced into a DhMonoFc domain (referred to herein as “DhMonoFc(N297G)”).

II.H.1 DhCpmFc(-)(N297G)-GDF15(Ndel3):DhCpmFc(+)(N297G)

The designation “DhCpmFc(-)(N297G)-GDF15(Ndel3):DhCpmFc(+)(N297G)” in the instant disclosure refers to a heterodimer comprising (i) a first polypeptide chain comprising a GDF15(Ndel3) polypeptide, the N-terminus of which is linked directly to the C-terminus of a DhCpmFc(-)(N297G) domain, and (ii) a second polypeptide chain comprising a DhCpmFc(+)(N297G) domain.

In certain embodiments, a tetramer is provided, comprising a dimer of two DhCpmFc(-)(N297G)-GDF15(Ndel3):DhCpmFc(+)(N297G) heterodimers in which the two first polypeptide chains of each heterodimer are linked via an interchain disulfide bond between their respective GDF15 regions.

More particularly, in a specific embodiment, the tetramer comprises:

(a) two DhCpmFc(+)(N297G) domains (one each heterodimer) comprising the sequence:

APELLGGPSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVD
 GVEVHNAKTKPREEQYGSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPA
 PIEKTISKAKGQPREPQVYTLPPSRKEMTKNQVSLTCLVKGFYPSDIAVE
 WESNGQPENNYKTTPPVLKSDGSFFLYSKLTVDKSRWQQGNVFSQVMHE
 5 ALHNHYTQKSLSLSPG (SEQ ID NO:287),

(b) two DhCpmFc(-)(N297G) domains (one each heterodimer) comprising the sequence:

APELLGGPSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVD
 GVEVHNAKTKPREEQYGSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPA
 10 PIEKTISKAKGQPREPQVYTLPPSREEMTKNQVSLTCLVKGFYPSDIAVE
 WESNGQPENNYDTTPPVLDSDGSFFLYSDLTVDKSRWQQGNVFSQVMHE
 ALHNHYTQKSLSLSPG (SEQ ID NO:132),

and

(c) two GDF15(Nde13) polypeptides (one each heterodimer) comprising the sequence
 15 of SEQ ID NO:55.

In a preferred embodiment, the first polypeptide chain comprises the amino acid sequence:

APELLGGPSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVD
 GVEVHNAKTKPREEQYGSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPA
 20 PIEKTISKAKGQPREPQVYTLPPSREEMTKNQVSLTCLVKGFYPSDIAVE
 WESNGQPENNYDTTPPVLDSDGSFFLYSDLTVDKSRWQQGNVFSQVMHE
 ALHNHYTQKSLSLSPGGDHCPGLGPRCCRLHTVRASLEDLGWADWVLSR
 EVQVTMCIGACPSQFRAANMHAQIKTSLHRLKPDTPAPCCVPASYNPMV
 LIQKTDGTGVSQTYYDDLLAKDCHCI (SEQ ID NO:134),

25 which is encoded by the nucleic acid sequence:

gcacctgaactcctggggggaccgtcagtccttcttccccccaaaacc
 caaggacaccctcatgatctcccggaacctgaggtcacatgcgtggtgg
 tggacgtgagccacgaagacctgaggtcaagttcaactggtacgtggac
 ggcgtggaggtgcataatgccaaagacaaagccgaggaggagcagtaagg
 30 gagcacgtaccgtgtggtcagcgtcctcaccgtcctgcaccaggactggc
 tgaatggcaaggagtacaagtgaaggtctccaacaaagccctcccagcc
 cccatcgagaaaaccatctccaaagccaaagggcagccccgagaaccaca
 ggtgtacaccctgcccccatcccgaggagatgaccaagaaccagggtca
 gcctgacctgcctggtcaaaggcttctatcccagcgacatcgccgtggag

tgggagagcaatgggcagccggagaacaactacgacaccacgcctcccgt
 gctggactccgacggctccttcttctctatagcgacctcaccgtggaca
 agagcaggtggcagcaggggaacgtcttctcatgctccgtgatgcatgag
 gctctgcacaaccactacacgcagaagagcctctccctgtctccgggtgg
 5 agaccactgtccgctcgggccccggcggttgetgccgtctgcacacggtec
 gcgcgtcgtggaagacctgggctgggcccattgggtgctgtcgccacgg
 gaggtgcaagtgacctgtgcatcggcgcgtgcccagaccagttccgggc
 ggcaaacatgcacgcgcagatcaagacgagcctgcaccgcctgaagcccg
 acacggtgccagcgccctgctgctgcccgccagctacaatcccatggtg
 10 ctcatcctcaaaagaccgacacccgggtgtcgtccagacctatgatgactt
 gttagccaaagactgccactgcata (SEQ ID NO:133) .

In a preferred embodiment, the second polypeptide chain comprises the amino acid sequence:

APELLGGPSVFLFPPKPKDTLMISRTPEVTCVVDVSHEDPEVKFNWYVD
 15 GVEVHNAKTKPREEQYGSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPA
 PIEKTISKAKGQPREPQVYTLPPSRKEMTKNQVSLTCLVKGFYPSDIAVE
 WESNGQPENNYKTTPPVLKSDGSFFLYSKLTVDKSRWQQGNVFSQVMHE
 ALHNHYTQKSLSLSPGK (SEQ ID NO:131) ,

which is encoded by the nucleic acid sequence:

20 gcacctgaactcctggggggaccgtcagtccttcttccccccaaaacc
 caaggacaccctcatgatctcccgaccctgaggtcacatgcgtgggtgg
 tggacgtgagccacgaagaccctgaggtcaagttcaactggtacgtggac
 ggcggtggaggtgcataatgccaaagacaaagccgcgggaggagcagtaagg
 gagcacgtaccgtgtggtcagcgtcctcaccgtcctgcaccaggactggc
 25 tgaatggcaaggagtacaagtgcaaggtctccaacaaagccctcccagcc
 cccatcgagaaaaccatctccaaagccaaagggcagccccgagaaccaca
 ggtgtacaccctgcccccatcccggaaggagatgaccaagaaccaggtca
 gcctgacctgcctggtcaaaggcttctatcccagcgacatcgccgtggag
 tgggagagcaatgggcagccggagaacaactacaagaccacgcctcccgt
 30 gctgaagtccgacggctccttcttctctatagcaagctcaccgtggaca
 agagcaggtggcagcaggggaacgtcttctcatgctccgtgatgcatgag
 gctctgcacaaccactacacgcagaagagcctctccctgtctccgggtaa
 atga (SEQ ID NO:135) .

As discussed above, a tetramer is provided comprising two polypeptide chains comprising the sequence of SEQ ID NO:134 and two polypeptide chains comprising the sequence of SEQ ID NO:131.

5 *II.H.2 DhCpmFc(-)(N297G)-GDF15(N3D):DhCpmFc(+)(N297G)*

The designation “DhCpmFc(-)(N297G)-GDF15(N3D):DhCpmFc(+)(N297G)” in the instant disclosure refers to a heterodimer comprising (i) a first polypeptide chain comprising a GDF15(N3D) sequence, the N-terminus of which is linked directly to the C-terminus of a DhCpmFc(-)(N297G) domain, and (ii) a second polypeptide chain comprising a
10 DhCpmFc(+)(N297G) domain.

In certain embodiments, a tetramer is provided, comprising a dimer of two DhCpmFc(-)(N297G)-GDF15(N3D):DhCpmFc(+)(N297G) heterodimers in which the two first polypeptide chains of each heterodimer are linked via an interchain disulfide bond between their respective GDF15 regions.

15 More particularly, in a specific embodiment, the tetramer comprises:

(a) two DhCpmFc(+)(N297G) domains (one each heterodimer) comprising the sequence of SEQ ID NO:287,

(b) two DhCpmFc(-)(N297G) domains (one each heterodimer) comprising the sequence of SEQ ID NO:132, and

20 (c) two GDF15(N3D) polypeptides (one each heterodimer) comprising the sequence of SEQ ID NO:52.

In a preferred embodiment, the first polypeptide chain comprises the amino acid sequence:

25 APELLGGPSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVD
GVEVHNAKTKPREEQYGSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPA
PIEKTISKAKGQPREPQVYTLPPSREEMTKNQVSLTCLVKGFYPSDIAVE
WESNGQPENNYDTTPPVLDSDGSFFLYSDLTVDKSRWQQGNVFSQSVME
ALHNHYTQKSLSLSPGARDGDHCPGPRCCRLHTVRASLEDLGWADWVL
SPREVQVTMCIGACPSQFRAANMHAQIKTSLHRLKPDTVPAPCCVPASYN
30 PMVLIQKTDGTGVSLSQTYDDLAKDCHCI (SEQ ID NO:137),

which is encoded by the nucleic acid sequence:

gcacctgaactcctggggggaccgtcagtccttcttccccccaaaacc
caaggacaccctcatgatctcccgaccctgaggtcacatgcgtggtgg
tggacgtgagccacgaagaccctgaggtcaagttcaactggtacgtggac
35 ggcgtggaggtgcataatgccaaagacaaagccgcggaggagcagtacgg

gagcacgtaccgtgtggtcagcgtcctcaccgtcctgcaccaggactggc
 tgaatggcaaggagtacaagtgaagggtctccaacaaagccctcccagcc
 cccatcgagaaaaaccatctccaaagccaaagggcagccccgagaaccaca
 ggtgtacaccctgcccccatccccgggaggagatgaccaagaaccaggtea
 5 gcttgacctgacctggtcaaaggcttctatcccagcgacatcgccgtggag
 tgggagagcaatgggcagccggagaacaactacgacaccacgcctcccgt
 gctggactccgacggctccttcttctctatagcgacctcaccgtggaca
 agagcaggtggcagcaggggaacgtcttctcatgctccgtgatgcatgag
 gctctgcacaaccactacacgcagaagagcctctccctgtctccgggtgc
 10 gcgcgacggagaccactgtccgctcgggccccgggcgttgctgccgtctgc
 acacggtccgcgctcgctggaagacctgggctgggccgattgggtgctg
 tcgccacgggaggtgcaagtgacctgtgcatcggcgctgcccagacca
 gttccgggcggaacatgcacgcgcagatcaagacgagcctgcaccgcc
 tgaagcccgacacggtgccagcgccctgctgctgcccgcagctacaat
 15 cccatggtgctcattcaaaagaccgacaccggggtgtcgctccagacct
 tgatgacttgtagccaaagactgccactgcata (SEQ ID NO:136).

In a preferred embodiment, the second polypeptide chain comprises the amino acid sequence of SEQ ID NO:131, which is encoded by the nucleic acid sequence of SEQ ID NO:135.

20 As discussed above, a tetramer is provided comprising two polypeptide chains comprising the sequence of SEQ ID NO:137 and two polypeptide chains comprising the sequence of SEQ ID NO:131.

II.H.3 *DhCpmFc(-)(N297G)-G₄-GDF15(N3D):DhCpmFc(+)(N297G)*

25 The designation “*DhCpmFc(-)(N297G)-G₄-GDF15(N3D):DhCpmFc(+)(N297G)*” in the instant disclosure refers to a heterodimer comprising (i) a first polypeptide chain comprising a GDF15(N3D) polypeptide linked to a *DhCpmFc(-)(N297G)* domain via a linker comprising the sequence of SEQ ID NO:58 that connects the N-terminus of the GDF15(N3D) polypeptide to the C-terminus of the *DhCpmFc(-)(N297G)* domain and (ii) a second
 30 polypeptide chain comprising a *DhCpmFc(+)(N297G)* domain.

In certain embodiments, a tetramer is provided, comprising a dimer of two *DhCpmFc(-)(N297G)-G₄-GDF15(N3D):DhCpmFc(+)(N297G)* heterodimers in which the two first polypeptide chains of each heterodimer are linked via an interchain disulfide bond between their respective GDF15 regions.

35 More particularly, in a specific embodiment, the tetramer comprises:

(a) two DhCpmFc(+)(N297G) domains (one each heterodimer) comprising the sequence of SEQ ID NO:287,

(b) two DhCpmFc(-)(N297G) domains (one each heterodimer) comprising the sequence of SEQ ID NO:132,

5 (c) two GDF15(N3D) polypeptides (one each heterodimer) comprising the sequence of SEQ ID NO:52, and

(d) two polypeptide linkers (one each heterodimer) comprising the sequence of SEQ ID NO:58 each linking the N-terminus of a GDF15(N3D) polypeptide to the C-terminus of a DhCpmFc(-)(N297G) domain via a peptide bond.

10 In a preferred embodiment, the first polypeptide chain comprises the amino acid sequence:

APELLGGPSVFLFPPKPKDTLMI SRTPEVTCVVVDVSHEDPEVKFNWYVD
GVEVHNAKTKPREEQYGSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPA
PIEKTISKAKGQPREPQVYTLPPSREEMTKNQVSLTCLVKGFYPSDIAVE
15 WESNGQPENNYDTPPVLDSDGSFFLYSDLTVDKSRWQQGNVFSCSVMHE
ALHNHYTQKSLSLSPGGGGGARDGDHCPLGPGRCRLHTVRASLEDLGWA
DWVLSPREVQVTMCIGACPSQFRAANMHAQIKTSLHRLKPDTPAPCCVP
ASYNPMVLIQKTDGTGVS LQTYDDLAKDCHCI (SEQ ID NO:139),

which is encoded by the nucleic acid sequence:

20 gcacctgaactcctggggggaccgtcagtccttcctcttccccccaaaacc
caaggacaccctcatgatctcccggaaccctgaggtcacatgcgtggtgg
tggacgtgagccacgaagaccctgaggtcaagttcaactggtacgtggac
ggcgtggaggtgcataatgccaaagacaaagccgcgggaggagcagtacgg
gagcacgtaccgtgtggtcagcgtcctcaccgtcctgcaccaggactggc
25 tgaatggcaaggagtacaagtgaaggtctccaacaaagccctcccagcc
cccatcgagaaaaccatctccaaagccaaagggcagccccgagaaccaca
ggtgtacaccctgcccccatcccgaggagatgaccaagaaccagggtca
gcctgacctgcctggtcaaaggcttctatcccagcgacatcgccgtggag
tgggagagcaatgggcagccggagaacaactacgacaccacgcctcccgt
30 gctggactccgacggctccttcttcctctatagcgacctcaccgtggaca
agagcaggtggcagcaggggaacgtcttctcatgctccgtgatgcatgag
gctctgcacaaccactacacgcagaagagcctctccctgtctccgggtgg
aggtggtggagcgcgcgacggagaccactgtccgctcgggcccggggcgtt
gctgccgtctgcacacgggtccgcgcgtcgctggaagacctgggctgggccc
35 gattgggtgctgtcgccacgggaggtgcaagtgacctatgtgcatcggcgc

gtgccccgagccagttccggggcggaacacatgcacgcgcagatcaagacga
 gcctgcaccgcctgaagcccgacacgggtgccagcgccctgctgcgtgccc
 gccagctacaatcccatgggtgctcattcaaaagaccgacaccgggggtgtc
 gctccagacctatgatgacttggttagccaaagactgccactgcata (SEQ ID

5 NO:138) .

In a preferred embodiment, the second polypeptide chain comprises the amino acid sequence of SEQ ID NO:131, which is encoded by the nucleic acid sequence of SEQ ID NO:135.

As discussed above, a tetramer is provided comprising two polypeptide chains comprising the sequence of SEQ ID NO:139 and two polypeptide chains comprising the sequence of SEQ ID NO:131.

II.H.4 DhCpmFc(-)(N297G)(Y349C)-GDF15(Ndel3):DhCpmFc(+)(N297G)(S354C)

The designation “DhCpmFc(-)(N297G)(Y349C)-GDF15(Ndel3):DhCpmFc(+)(N297G)(S354C)” in the instant disclosure refers to a heterodimer comprising (i) a first polypeptide chain comprising a GDF15(Ndel3) polypeptide, the N-terminus of which is linked directly to the C-terminus of a DhCpmFc(-)(N297G)(Y349C) domain, and (ii) a second polypeptide chain comprising a DhCpmFc(+)(N297G)(S354C) domain. The cysteine clamp mutations allow the first and second polypeptide chains to be linked via an interchain disulfide bond between C349 of the first polypeptide chain and C354 of the second polypeptide chain.

In certain embodiments, a tetramer is provided, comprising a dimer of two DhCpmFc(-)(N297G)(Y349C)-GDF15(Ndel3):DhCpmFc(+)(N297G)(S354C) heterodimers in which the two first polypeptide chains of each heterodimer are linked via an interchain disulfide bond between their respective GDF15 regions.

More particularly, in a specific embodiment, the tetramer comprises:

(a) two DhCpmFc(+)(N297G)(S354C) domains (one each heterodimer) comprising the sequence:

APELLGGPSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVD
 GVEVHNAKTKPREEQYGSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPA
 PIEKTISKAKGQPREPQVYTLPPCRKEMTKNQVSLTCLVKGFYPSDIAVE
 WESNGQPENNYKTTPPVLKSDGSFFLYSKLTVDKSRWQQGNVFSQVMHE
 ALHNYHTQKSLSLSPG (SEQ ID NO:288) ,

(b) two DhCpmFc(-)(N297G)(Y349C) domains (one each heterodimer) comprising the sequence:

APELLGGPSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVD
 GVEVHNAKTKPREEQYGSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPA
 PIEKTISKAKGQPREPQVCTLPSSREEMTKNQVSLTCLVKGFYPSDIAVE
 WESNGQPENNYDTTPPVLDSDGSFFLYSDLTVDKSRWQQGNVFSQSVME

5 ALHNHYTQKSLSLSPG (SEQ ID NO:141),

and

(c) two GDF15(Nde13) polypeptides (one each heterodimer) comprising the sequence of SEQ ID NO:55.

In a preferred embodiment, the first polypeptide chain comprises the amino acid
 10 sequence:

APELLGGPSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVD
 GVEVHNAKTKPREEQYGSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPA
 PIEKTISKAKGQPREPQVCTLPSSREEMTKNQVSLTCLVKGFYPSDIAVE
 WESNGQPENNYDTTPPVLDSDGSFFLYSDLTVDKSRWQQGNVFSQSVME
 15 ALHNHYTQKSLSLSPGGDHCPGPRCCRLHTVRASLEDLGWADWVLSR
 EVQVTMCIGACPSQFRAANMHAQIKTSLHRLKPDTPAPCCVPASYNPMV
 LIQKTDGTGSLQTYDDLAKDCHCI (SEQ ID NO:143),

which is encoded by the nucleic acid sequence:

gcacctgaactcctggggggaccgtcagtcctcctcttccccccaaaacc
 20 caaggacaccctcatgatctcccgaccctgaggtcacatgcgtggtgg
 tggacgtgagccacgaagaccctgaggtcaagttcaactggtacgtggac
 ggcgtggaggtgcataatgccaaagacaaagccgaggagcagtacgg
 cagcacgtaccgtgtggtcagcgtcctcaccgtcctgcaccaggactggc
 tgaatggcaaggagtacaagtgaaggtctccaacaaagccctcccagcc
 25 cccatcgagaaaaccatctccaaagccaaagggcagccccgagaaccaca
 ggtgtgcaccctgcccccatcccgaggagatgaccaagaaccaggtca
 gcctgacctgcctggtcaaaggcttctatcccagcgacatcgccgtggag
 tgggagagcaatgggcagccggagaacaactacgacaccacgcctcccgt
 gctggactccgacggtccttcttctctatagcgacctcaccgtggaca
 30 agagcaggtggcagcaggggaacgtcttctcatgctccgtgatgcatgag
 gctctgcacaaccactacacgcagaagagcctctccctgtctccgggtgg
 agaccactgtccgctcgggcccggcggttgctgccgtctgcacacgggtcc
 gcgcgtcgctggaagacctgggctgggcccattgggtgctgtcgccacgg
 gaggtgcaagtgacctgtgcatcggcgcgtgcccagaccagttccgggc
 35 ggcaaacatgcacgcgcatcaagacgagcctgcaccgcctgaagcccg

acacggtgccagcgccctgctgctgccccgccagctacaatcccatggtg
ctcattcaaaagaccgacacccgggtgctgctccagacctatgatgactt
gttagccaaagactgccactgcata (SEQ ID NO:142).

In a preferred embodiment, the second polypeptide chain comprises the amino acid
5 sequence:

APELLGGPSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVD
GVEVHNAKTKPREEQYGSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPA
PIEKTISKAKGQPREPQVYTLPPCRKEMTKNQVSLTCLVKGFYPSDIAVE
WESNGQPENNYKTTTPVLKSDGSFFLYSKLTVDKSRWQQGNVFSCSVMHE
10 ALHNHYTQKSLSLSPGK (SEQ ID NO:140),

which is encoded by the nucleic acid sequence:

gccccagagctgcttggtggaccatccgtgttcctgtttcctccaaagcc
gaaggacaccctgatgatctcaagaactccggaagtgacttgctgctgctg
tggacgtgtcacatgaggatccagaggtcaagttcaattggtatgtggac
15 ggagtggaagtgcataacgcccaagaccaaaccggaagaacagtacgg
gagcacctaccgctggtgagcgtccttactgtgctccaccaggactggc
ttaatgggaaggaatacaagtgttaagggtgtccaacaaggccctccccgct
cccatcgaaaagaccatctcaaaggcaaaggggcaaccaagggaacctca
agtgtacaccctgcctccgtgcaggaaggagatgaccaagaaccagggtca
20 gcctgacttgctcgtgaagggttctatcccagcgatattgctgtggaa
tgggagtgcaaatggccagcccagagaataactacaaaactaccccaccgct
gctgaaatctgatgggtccttcttcttcttactccaagctgaccgtggaca
agagccgctggcaacaaggcaatgtctttagctgctcagtgatgcattgag
gctctccataatcactacactcagaagtcactgtccctgtctccgggttaa
25 a (SEQ ID NO:144).

As discussed above, a tetramer is provided comprising two polypeptide chains
comprising the sequence of SEQ ID NO:143 and two polypeptide chains comprising the
sequence of SEQ ID NO:140.

30 II.H.5 *DhCpmFc(-)(N297G)(Y349C)-GDF15(N3D):DhCpmFc(+)(N297G)(S354C)*

The designation “DhCpmFc(-)(N297G)(Y349C)-
GDF15(N3D):DhCpmFc(+)(N297G)(S354C)” in the instant disclosure refers to a
heterodimer comprising (i) a first polypeptide chain comprising a GDF15(N3D) polypeptide,
the N-terminus of which is linked directly to the C-terminus of a

DhCpmFc(-)(N297G)(Y349C) domain, and (ii) a second polypeptide chain comprising a DhCpmFc(+)(N297G)(S345C) domain, The cysteine clamp mutations allow the first and second polypeptide chains to be linked via an interchain disulfide bond between C349 of the first polypeptide chain and C354 of the second polypeptide chain.

- 5 In certain embodiments, a tetramer is provided, comprising a dimer of two DhCpmFc(-)(N297G)(Y349C)-GDF15(N3D):DhCpmFc(+)(N297G)(S354C) heterodimers in which the two first polypeptide chains of each heterodimer are linked via an interchain disulfide bond between their respective GDF15 regions.

More particularly, in a specific embodiment, the tetramer comprises:

- 10 (a) two DhCpmFc(+)(N297G)(S354C) domains (one each heterodimer) comprising the sequence of SEQ ID NO:288,
 (b) two DhCpmFc(-)(N297G)(Y349C) domains comprising the sequence of SEQ ID NO:141, and
 (c) two GDF15(N3D) polypeptides (one each heterodimer) comprising the sequence
 15 of SEQ ID NO:52.

In a preferred embodiment, the first polypeptide chain comprises the amino acid sequence:

APELLGGPSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVD
 GVEVHNAKTKPREEQYGSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPA
 20 PIEKTISKAKGQPREPQVCTLPSSREEMTKNQVSLTCLVKGFYPSDIAVE
 WESNGQPENNYDTPPVLDSDGSFFLYSDLTVDKSRWQQGNVFSCSVMHE
 ALHNYHTQKSLSLSPGARDGDHCPLGPGRCRLHTVRASLEDLGWADWVL
 SPREVQVTMCIGACPSQFRAANMHAQIKTSLHRLKPDTPAPCCVPASYN
 PMVLIQKTDGTGVSLSQTYDDLAKDCHCI (SEQ ID NO:146),

- 25 which is encoded by the nucleic acid sequence:

gcacctgaactcctggggggaccgctcagtccttcctcttccccccaaaacc
 caaggacaccctcatgatctcccgaccctgaggtcacatgcgtggtgg
 tggacgtgagccacgaagaccctgaggtcaagttcaactggtacgtggac
 ggcgtggaggtgcataatgccaaagacaaagccgcgggaggagcagtacgg
 30 gagcacgtaccgtgtggtcagcgtcctcaccgtcctgcaccaggactggc
 tgaatggcaaggagtacaagtgaaggtctccaacaaagccctcccagcc
 cccatcgagaaaaccatctccaaagccaaagggcagccccgagaaccaca
 ggtgtgcaccctgcccccatcccgaggagatgaccaagaaccaggtca
 gcctgacctgcctggtcaaaggcttctatcccagcgacatcgccgtggag
 35 tgggagagcaatgggcagccggagaacaactacgacaccacgcctcccg

gctggactccgacggctccttcttctctatagcgacctcacctgggaca
agagcaggtggcagcaggggaacgtcttctcatgctccgtgatgcatgag
gctctgcacaaccactacacgcagaagagcctctccctgtctccgggtgc
gcgcgacggagaccactgtccgctcgggcccgggcggttgctgccgtctgc
5 acacgggtccgcgcgtcgctggaagacctgggctgggcccattgggtgctg
tcgccacgggaggtgcaagtgaccatgtgcatcggcgcgtgcccagagcca
gttccgggcggaacatgcacgcgcagatcaagacgagcctgcaccgcc
tgaagcccgacacggtgccagcgcctgctgcgtgcccgcagctacaat
cccatggtgctcattcaaaagaccgacaccggggtgctcgtccagacct
10 tgatgacttgtagccaaagactgccactgcata (SEQ ID NO:145).

In a preferred embodiment, the second polypeptide chain comprises the amino acid sequence of SEQ ID NO:140, which is encoded by the nucleic acid sequence of SEQ ID NO:144.

As discussed above, a tetramer is provided comprising two polypeptide chains comprising the sequence of SEQ ID NO:146 and two polypeptide chains comprising the sequence of SEQ ID NO:140.

II.H.6 DhCpmFc(-)(N297G)(L351C)-GDF15(Ndel3):DhCpmFc(+)(N297G)(L351C)

The designation “DhCpmFc(-)(N297G)(L351C)-GDF15(Ndel3):DhCpmFc(+)(N297G)(L351C)” in the instant disclosure refers to a heterodimer comprising (i) a first polypeptide chain comprising a GDF15(Ndel3) polypeptide, the N-terminus of which is linked directly to the C-terminus of a DhCpmFc(-)(N297G)(L351C) domain, and (ii) a second polypeptide chain comprising a DhCpmFc(+)(N297G)(L351C) domain. The cysteine clamp mutations allow the first and second polypeptide chains to be linked via an interchain disulfide bond between C351 of the first polypeptide chain and C351 of the second polypeptide chain.

In certain embodiments, a tetramer is provided, comprising a dimer of two DhCpmFc(-)(N297G)(L351C)-GDF15(Ndel3):DhCpmFc(+)(N297G)(L351C) heterodimers in which the two first polypeptide chains of each heterodimer are linked via an interchain disulfide bond between their respective GDF15 regions.

More particularly, in a specific embodiment, the tetramer comprises:

(a) two DhCpmFc(+)(N297G)(L351C) domains (one each heterodimer) comprising the sequence:

APELLGGPSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVD
35 GVEVHNAKTKPREEQYGSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPA
PIEKTISKAKGQPREPQVYTCPPSREEMTKNQVSLTCLVKGFYPSDIAVE

WESNGQPENNYKTTPPVLSKSDGSFFLYSKLTVDKSRWQQGNVFSCSVME
 ALHNHYTQKSLSLSPG (SEQ ID NO:289),

(b) two DhCpmFc(-)(N297G)(L351C) domains (one each heterodimer) comprising the sequence:

5 APELLGGPSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVD
 GVEVHNAKTKPREEQYGSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPA
 PIEKTISKAKGQPREPQVYTCPPSREEMTKNQVSLTCLVKGFYPSDIAVE
 WESNGQPENNYDTTPPVLDSDGSFFLYSDLTVDKSRWQQGNVFSCSVME
 ALHNHYTQKSLSLSPG (SEQ ID NO:148),

10 and

(c) two GDF15(Nde13) polypeptides (one each heterodimer) comprising the sequence of SEQ ID NO:55.

In a preferred embodiment, the first polypeptide chain comprises the amino acid sequence:

15 APELLGGPSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVD
 GVEVHNAKTKPREEQYGSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPA
 PIEKTISKAKGQPREPQVYTCPPSRKEMTKNQVSLTCLVKGFYPSDIAVE
 WESNGQPENNYKTTPPVLSKSDGSFFLYSKLTVDKSRWQQGNVFSCSVME
 ALHNHYTQKSLSLSPGARNGDHCPLGPGRCCRLHTVRASLEDLGWADWVL
 20 SPREVQVTMCIGACPSQFRAANMHAQIKTSLHRLKPDTPAPCCVPASYN
 PMVLIQKTDGTGVSLLQTYDDLAKDCHCI (SEQ ID NO:150),

which is encoded by the nucleic acid sequence:

gcgcgcgaactgctggcgccgagcgtgtttctgtttccgccgaaacc
 gaaagataccctgatgattagccgcaccccggaagtgcctgctggtgg
 25 tggatgtgagccatgaagatccggaagtgaatattaactggtatgtggat
 ggcgtggaagtgcataacgcgaaaaccaaaccgcgcgaagaacagtatgg
 cagcacctatcgctggtgagcgtgctgaccgtgctgcatcaggattggc
 tgaacggcaaaagaatataaatgcaaagtgagcaaaaagcgtgccggcg
 ccgattgaaaaaaccattagcaaagcgaaaggccagccgcgcgaaccgca
 30 ggtgtatacctgcccgcgagccgcaaagaaatgacaaaaaccaggtga
 gcctgacctgcctggtgaaaggcttttatccgagcgatattgcggtggaa
 tgggaaagcaacggccagccgaaaacaactataaaaccaccccgccggt
 gctgaaaagcgatggcagctttttctgtatagcaaactgaccgtggata
 aaagccgctggcagcagggaacgtgttagctgcagcgtgatgcatgaa

gcgctgcataaccattataccagaaaaagcctgagcctgagcccgggcgc
 gcgcaacggcgatcattgcccgctgggcccggcgctgctgccgcctgc
 ataccgtgcgcgcgagcctggaagatctgggctggcggttgggtgctg
 agcccgcgcaagtgcaggtgaccatgtgcattggcgctgcccgagcca
 5 gtttcgcgcggcgaacatgcatgcgcagattaaaaccagcctgcacgcc
 tgaaaccggataccgtgccggcgccgtgctgcgtgccggcgagctataac
 ccgatggtgctgattcagaaaaccgataccggcgtgagcctgcagaccta
 tgatgatctgctggcgaaagattgccattgcatt (SEQ ID NO:149).

In a preferred embodiment, the second polypeptide chain comprises the amino acid
 10 sequence:

APELLGGPSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVD
 GVEVHNAKTKPREEQYGSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPA
 PIEKTISKAKGQPREPQVYTCPPSREEMTKNQVSLTCLVKGFYPSDIAVE
 WESNGQPENNYKTTTPVLKSDGSFFLYSKLTVDKSRWQQGNVVFSCSVMHE
 15 ALHNHYTQKSLSLSPGK (SEQ ID NO:147),

which is encoded by the nucleic acid sequence:

gcacctgaactcctggggggaccgtcagtccttcctcttccccccaaaacc
 caaggacaccctcatgatctcccgaccctgaggtcacatgcgtggtgg
 tggacgtgagccacgaagaccctgaggtcaagttcaactggtacgtggac
 20 ggcgtggaggtgcataatgccaaagacaaagccgcgggaggagcagtacgg
 gagcacgtaccgtgtggtcagcgtcctcaccgtcctgcaccaggactggc
 tgaatggcaaggagtacaagtgcaaggtctccaacaaagccctcccagcc
 cccatcgagaaaaccatctccaaagccaaagggcagccccgagaaccaca
 ggtgtacacctgtcccccatccgggaggagatgaccaagaaccaggtca
 25 gcctgacctgcctggtcaaaggcttctatcccagcgacatcgccgtggag
 tgggagagcaatgggcagccggagaacaactacgacaccacgcctcccgt
 gctggactccgacggtccttcttctctatagcgacctcaccgtggaca
 agagcaggtggcagcaggggaacgtcttctcatgctccgtgatgcatgag
 gctctgcacaaccactacacgcagaagagcctctccctgtctccgggtaa
 30 a (SEQ ID NO:151).

As discussed above, a tetramer is provided comprising two polypeptide chains
 comprising the sequence of SEQ ID NO:150 and two polypeptide chains comprising the
 sequence of SEQ ID NO:147.

II.H.7 DhCpmFc(-)(N297G)(L351C)-GDF15(N3D):DhCpmFc(+)(N297G)(L351C)

The designation “DhCpmFc(-)(N297G)(L351C)-GDF15(N3D):DhCpmFc(+)(N297G)(L351C)” in the instant disclosure refers to a heterodimer comprising (i) a first polypeptide chain comprising a GDF15(N3D) polypeptide, the N-terminus of which is linked directly to the C-terminus of a DhCpmFc(-)(N297G)(L351C) domain by a peptide bond and (ii) a second polypeptide chain comprising a DhCpmFc(+)(N297G)(L351C) domain. The cysteine clamp mutations allow the first and second polypeptide chains to be linked via an interchain disulfide bond between C351 of the first polypeptide chain and C351 of the second polypeptide chain.

In certain embodiments, a tetramer is provided, comprising a dimer of two DhCpmFc(-)(N297G)(L351C)-GDF15(N3D):DhCpmFc(+)(N297G)(L351C) heterodimers in which the two first polypeptide chains of each heterodimer are linked via an interchain disulfide bond between their respective GDF15 regions.

More particularly, in a specific embodiment, the tetramer comprises:

(a) two DhCpmFc(+)(N297G)(L351C) domains (one each heterodimer) comprising the sequence of SEQ ID NO:289,

(b) two DhCpmFc(-)(N297G)(L351C) domains (one each heterodimer) comprising the sequence of SEQ ID NO:148, and

(c) two GDF15(N3D) polypeptides (one each heterodimer) comprising the sequence of SEQ ID NO:52.

In a preferred embodiment, the first polypeptide chain comprises the amino acid sequence:

APELLGGPSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVD
GVEVHNAKTKPREEQYGSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPA
PIEKTISKAKGQPREPQVYTCPPSRKEMTKNQVSLTCLVKGFYPSDIAVE
WESNGQPENNYKTTTPVLKSDGSFFLYSKLTVDKSRWQQGNVFSQSVME
ALHNHYTQKSLSLSPGARDGDHCPLGPGRCRLHTVRASLEDLGWADWVL
SPREVQVTMCIGACPSQFRAANMHAQIKTSLHRLKPDTVPAPCCVPASYN
PMVLIQKTDGTGVSLQTYDDLLAKDCHCI (SEQ ID NO:153),

which is encoded by the nucleic acid sequence:

gcgccggaactgctggcgccgcccagcgctgtttctgtttccgccgaaacc
gaaagataccctgatgattagccgcaccccggaagtgcctgcgtggtgg
tggatgtgagccatgaagatccggaagtgaaatttaactggtatgtggat
ggcgtggaagtgcataacgcgaaaaccaaaccgcgcgaagaacagtatgg
cagcacctatcgcggtggtgagcgtgctgaccgtgctgcatcaggattggc

tgaacggcacaagaatataaatgcaaagtgagcaacaaagcgctgccggcg
 ccgattgaaaaaaccattagcaaagcgaaaggccagccgcggaaccgca
 ggtgtatacctgcccgcgagccgcaaagaaatgacaaaaaccaggtga
 gcctgacctgcctggtgaaaggcttttatccgagcgatattgcggtggaa
 5 tgggaaagcaacggccagccggaaaacaactataaaaccaccccgccggt
 gctgaaaagcgatggcagcttttttctgtatagcaaactgaccgtggata
 aaagccgctggcagcagggcaacgtgttttagctgcagcgtgatgcatgaa
 gcgctgcataaccattataccagaaaagcctgagcctgagcccgggcgc
 gcgcgatggcgatcattgcccgctgggcccggccgctgctgccgcctgc
 10 ataccgtgcgcgcgagcctggaagatctgggctgggaggattgggtgctg
 agcccgcgcaagtgcaggtgaccatgtgcattggcgcgtgcccgagcca
 gtttcgcgcggcgaacatgcatgcgcagattaaaaccagcctgcacgcc
 tgaaaccggataccgtgccggcgccgtgctgcgtgccggcgagctataac
 ccgatgggtgctgattcagaaaaccgataccggcgctgagcctgcagaccta
 15 tgatgatctgctggcgaaagattgccattgcatt (SEQ ID NO:152) .

In a preferred embodiment, the second polypeptide chain comprises the amino acid sequence of SEQ ID NO:147, which is encoded by the nucleic acid sequence of SEQ ID NO:151.

As discussed above, a tetramer is provided comprising two polypeptide chains
 20 comprising the sequence of SEQ ID NO:153 and two polypeptide chains comprising the
 sequence of SEQ ID NO:147.

II.H.8 DhCpmFc(-)(N297G)(A287C)-GDF15(Ndel3):DhCpmFc(+)(N297G)(L306C)

The designation “DhCpmFc(-)(N297G)(A287C)-
 25 GDF15(Ndel3):DhCpmFc(+)(N297G)(L306C)” in the instant disclosure refers to a
 heterodimer comprising (i) a first polypeptide chain comprising a GDF15(Ndel3) polypeptide,
 the N-terminus of which is linked directly to the C-terminus of a
 DhCpmFc(-)(N297G)(A287C) domain, and (ii) a second polypeptide chain comprising a
 DhCpmFc(+)(N297G)(L306C) domain.

30 In certain embodiments, a tetramer is provided, comprising a dimer of two
 DhCpmFc(-)(N297G)(A287C)-GDF15(Ndel3):DhCpmFc(+)(N297G)(L306C) heterodimers
 in which the two first polypeptide chains of each heterodimer are linked via an interchain
 disulfide bond between their respective GDF15 regions.

More particularly, in a specific embodiment, the tetramer comprises:

35 (a) two DhCpmFc(+)(N297G)(L306C) domains (one each heterodimer) comprising
 the sequence:

APELLGGPSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVD
 GVEVHNCKTKPREEQYGSTYRVVSVCTV LHQDWLNGKEYKCKVSNKALPA
 PIEKTISKAKGQPREPQVYTLPPSRKEMTKNQVSLTCLVKGFYPSDIAVE
 WESNGQPENNYKTTPPV LKSDGSFFLYSKLTVDKSRWQQGNV FSCSVMHE
 5 ALHNHYTQKSLSLSPG (SEQ ID NO:290),

(b) two DhCpmFc(-)(N297G)(A287C) domains (one each heterodimer) comprising the sequence:

APELLGGPSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVD
 GVEVHNCKTKPREEQYGSTYRVVSVCTV LHQDWLNGKEYKCKVSNKALPA
 10 PIEKTISKAKGQPREPQVYTLPPSREEMTKNQVSLTCLVKGFYPSDIAVE
 WESNGQPENNYDTTPPV LKSDGSFFLYSDLTVDKSRWQQGNV FSCSVMHE
 ALHNHYTQKSLSLSPG (SEQ ID NO:268),

and

(c) two GDF15(Nde13) polypeptides (one each heterodimer) comprising the sequence
 15 of SEQ ID NO:55.

In a preferred embodiment, the first polypeptide chain comprises the amino acid sequence:

APELLGGPSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVD
 GVEVHNCKTKPREEQYGSTYRVVSVCTV LHQDWLNGKEYKCKVSNKALPA
 20 PIEKTISKAKGQPREPQVYTLPPSREEMTKNQVSLTCLVKGFYPSDIAVE
 WESNGQPENNYDTTPPV LKSDGSFFLYSDLTVDKSRWQQGNV FSCSVMHE
 ALHNHYTQKSLSLSPGGDHCP LGPGRCCRLHTVRASLEDLGWADWVLSR
 EVQVTMCIGACPSQFRAANMHAQIKTSLHRLKPDTPAPCCVPASYNPMV
 LIQKTD TGVS LQTYDDLAKDCHCI (SEQ ID NO:269),

25 which is encoded by the nucleic acid sequence:

gcacctgaactcctggggggaccgtcagtccttctcttcccc
 caaaacccaaggacacctcatgatctcccggaacctgaggtcacatgc
 gtggtggtggacgtgagccacgaagacctgaggtcaagttcaactggt
 cgtggacggcgtggaggtgcataattgcaagacaaagccgcgggaggagc
 30 agtacggcagcacgtaccgtgtggtcagcgtctgcaccgtcctgcaccag
 gactggctgaatggcaaggagtacaagtgaaggtctccaacaaagccct
 cccagcccccatcgagaaaaccatctccaaagccaaagggcagccccgag
 aaccacaggtgtacacctgcccccatcccgaggagatgaccaagaac
 caggtcagcctgacctgctggtcaaaggcttctatcccagcgacatcgc

cgtggagtgaggagcaatgggcagccggagaacaactacgacaccacgc
 ctcccggtgctggactccgacggctccttcttctctatagcgacctcacc
 gtggacaagagcaggtggcagcaggggaacgtcttctcatgctccgtgat
 gcatgaggctctgcacaaccactacacgcagaagagcctctccctgtctc
 5 cgggtggagaccactgtccgctcgggcccggcggttgctgccgtctgcac
 acggtccgcgcgtcgtggaagacctgggctgggccgattgggtgctgtc
 gccacgggaggtgcaagtgacctgtgcatcggcgcgtgcccagaccagt
 tccgggcggaacatgcacgcgcagatcaagacgagcctgcaccgcctg
 aagcccgacacggtgccagcgccctgctgctgcccgcagctacaatcc
 10 catggtgctcattcaaaagaccgacaccggggtgtcgtccagacctatg
 atgacttgtagccaaagactgccactgcata (SEQ ID NO:270).

In a preferred embodiment, the second polypeptide chain comprises the amino acid sequence:

APELLGGPSVFLFPPKPKDTLMISRTPEVTCVVDVSHEDPEVKFNWYVD
 15 GVEVHNCKTKPREEQYGSTYRVSVCTVLHQDWLNGKEYKCKVSNKALPA
 PIEKTISKAKGQPREPQVYTLPPSRKEMTKNQVSLTCLVKGFYPSDIAVE
 WESNGQPENNYKTTPPVLKSDGSFFLYSKLTVDKSRWQQGNVFSQVMHE
 ALHNHYTQKSLSLSPGK (SEQ ID NO:267),

which is encoded by the nucleic acid sequence:

20 gcacctgaactcctggggggaccgtcagtccttctcttcccc
 caaaacccaaggacacctcatgatctcccgaccctgaggtcacatgc
 gtggtggtggacgtgagccacgaagacctgaggtcaagttcaactggta
 cgtggacggcgtggaggtgcataattgcaagacaaagccgcgggaggagc
 agtacggcagcacgtaccgtgtggtcagcgtctgcaccgtcctgcaccag
 25 gactggctgaatggcaaggagtacaagtgaaggtctccaacaaagccct
 cccagcccccatcgagaaaaccatctccaaagccaaagggcagccccgag
 aaccacaggtgtacacctgcccccatcccggaaggagatgaccaagaac
 caggtcagcctgacctgcttgggtcaaaggcttctatcccagcgacatcgc
 cgtggagtgaggagcaatgggcagccggagaacaactacaagaccacgc
 30 ctcccggtgctgaagtccgacggctccttcttctctatagcaagctcacc
 gtggacaagagcaggtggcagcaggggaacgtcttctcatgctccgtgat
 gcatgaggctctgcacaaccactacacgcagaagagcctctccctgtctc
 cgggtaaa (SEQ ID NO:271).

As discussed above, a tetramer is provided comprising two polypeptide chains comprising the sequence of SEQ ID NO:269 and two polypeptide chains comprising the sequence of SEQ ID NO:267.

5 *II.H.9 DhCpmFc(-)(N297G)(A287C)-GDF15(N3D):DhCpmFc(+)(N297G)(L306C)*

The designation “DhCpmFc(-)(N297G)(A287C)-GDF15(N3D):DhCpmFc(+)(N297G)(L306C)” in the instant disclosure refers to a heterodimer comprising (i) a first polypeptide chain comprising a GDF15(N3D) polypeptide, the N-terminus of which is linked directly to the C-terminus of a
10 DhCpmFc(-)(N297G)(A287C) domain, and (ii) a second polypeptide chain comprising a DhCpmFc(+)(N297G)(L306C) domain.

In certain embodiments, a tetramer is provided, comprising a dimer of two DhCpmFc(-)(N297G)(A287C)-GDF15(N3D):DhCpmFc(+)(N297G)(L306C) heterodimers in which the two first polypeptide chains of each heterodimer are linked via an interchain
15 disulfide bond between their respective GDF15 regions.

More particularly, in a specific embodiment, the tetramer comprises:

- (a) two DhCpmFc(+)(N297G)(L306C) domains (one each heterodimer) comprising the sequence of SEQ ID NO:290,
- (b) two DhCpmFc(-)(N297G)(A287C) domains (one each heterodimer) comprising
20 the sequence of SEQ ID NO:268,
- and
- (c) two GDF15(N3D) polypeptides (one each heterodimer) comprising the sequence of SEQ ID NO:52.

In a preferred embodiment, the first polypeptide chain comprises the amino acid
25 sequence:

APELLGGPSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVD
GVEVHNCKTKPREEQYGSTYRVVSVCTV LHQDWLNGKEYKCKVSNKALPA
PIEKTISKAKGQPREPQVYTLPPSREEMTKNQVSLTCLVKGFYPSDIAVE
WESNGQPENNYDTTPPVLDSDGSFFLYSDLTVDKSRWQQGNV FSCSV MHE
30 ALHNHYTQKSLSLSPGARDGDHCPLGPGRCCRLHTVRASLEDLGWADWVL
SPREVQVTMCIGACPSQFRAANMHAQIKTSLHRLKPDTPAPCCVPASYN
PMVLIQKTDGTGVS LQTYDDLAKDCHCI (SEQ ID NO:272),

which is encoded by the nucleic acid sequence:

gcacctgaactcctggggggaccgtcagtccttctcttcccc
35 caaaacccaaggacaccctcatgatctcccgaccctgaggtcacatgc

gtggtggtggacgtgagccacgaagaccctgaggtcaagttcaactggta
 cgtggacggcgtggaggtgcataattgcaagacaaagccgcgaggagc
 agtacggcagcacgtaccgtgtggtcagcgtctgcaccgtcctgcaccag
 gactggctgaatggcaaggagtacaagtgcaaggtctccaacaaagccct
 5 cccagcccccatcgagaaaaccatctccaaagccaaagggcagccccgag
 aaccacaggtgtacaccctgcccccatcccgaggagatgaccaagaac
 caggtcagcctgacctgcctgggtcaaaggcttctatcccagcgacatcgc
 cgtggagtgaggagcaatgggcagccggagaaactacgacaccacgc
 ctcccggtgctggactccgacggctccttcttctctatagcgacctcacc
 10 gtggacaagagcaggtggcagcaggggaacgtcttctcatgctccgtgat
 gcatgaggctctgcacaaccactacacgcagaagagcctctccctgtctc
 cgggtgcgcgcgacggagaccactgtccgctcgggcccggggttgcgc
 cgtctgcacacgggtccgcgcgtcgtggaagacctgggctgggcccattg
 ggtgctgtcgccacgggaggtgcaagtgacctgtgcatcggcgcgtgcc
 15 cgagccagttccgggcggaacatgcacgcgcagatcaagacgagcctg
 caccgcctgaagcccgcacgggtgccagcgccctgctgctgcccgcag
 ctacaatcccatgggtgctcattcaaaagaccgacaccggggtgctgctcc
 agacctatgatgacttgtagccaaagactgccactgcata (SEQ ID NO:273) .

In a preferred embodiment, the second polypeptide chain comprises the amino acid
 20 sequence of SEQ ID NO:267, which is encoded by the nucleic acid sequence of SEQ ID
 NO:271.

As discussed above, a tetramer is provided comprising two polypeptide chains
 comprising the sequence of SEQ ID NO:272 and two polypeptide chains comprising the
 sequence of SEQ ID NO:267.

25

*II.H.10 DhCpmFc(-)(N297G)(A287C)(Y349C)-GDF15(Ndel3):
 DhCpmFc(+)(N297G)(L306C)(S354C)*

The designation “DhCpmFc(-)(N297G)(A287C)(Y349C)-
 30 GDF15(Ndel3):DhCpmFc(+)(N297G)(L306C)(S354C)” in the instant disclosure refers to a
 heterodimer comprising (i) a first polypeptide chain comprising a GDF15(Ndel3) polypeptide,
 the N-terminus of which is linked directly to the C-terminus of a
 DhCpmFc(-)(N297G)(A287C)(Y349C) domain, and (ii) a second polypeptide chain
 comprising a DhCpmFc(+)(N297G)(L306C)(S354C) domain.

35 In certain embodiments, a tetramer is provided, comprising a dimer of two
 DhCpmFc(-)(N297G)(A287C)(Y349C)-
 GDF15(Ndel3):DhCpmFc(+)(N297G)(L306C)(S354C) heterodimers in which the two first

polypeptide chains of each heterodimer are linked via an interchain disulfide bond between their respective GDF15 regions.

More particularly, in a specific embodiment, the tetramer comprises:

- (a) two DhCpmFc(+)(N297G)(L306C)(Y349C) domains (one each heterodimer) comprising the sequence:

APELLGGPSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVD
GVEVHNCKTKPREEQYGSTYRVVSVCTV LHQDWLNGKEYKCKVSNKALPA
PIEKTISKAKGQPREPQVYTLPPCRKEMTKNQVSLTCLVKGFYPSDIAVE
WESNGQPENNYKTTPPV LKSDGSFFLYSKLTVDKSRWQQGNV FSCSV MHE
10 ALHNHYTQKSLSLSPG (SEQ ID NO:291),

(b) two DhCpmFc(-)(N297G)(A287C)(S354C) domains (one each heterodimer) comprising the sequence:

APELLGGPSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVD
GVEVHNCKTKPREEQYGSTYRVVSVCTV LHQDWLNGKEYKCKVSNKALPA
15 PIEKTISKAKGQPREPQVCTLPSSREEMTKNQVSLTCLVKGFYPSDIAVE
WESNGQPENNYDTTPPV LDDSDGSFFLYSDLTVDKSRWQQGNV FSCSV MHE
ALHNHYTQKSLSLSPG (SEQ ID NO:275),

and

- (c) two GDF15(Ndel3) polypeptides (one each heterodimer) comprising the sequence of SEQ ID NO:55.

In a preferred embodiment, the first polypeptide chain comprises the amino acid sequence:

APELLGGPSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVD
GVEVHNCKTKPREEQYGSTYRVVSVCTV LHQDWLNGKEYKCKVSNKALPA
25 PIEKTISKAKGQPREPQVCTLPSSREEMTKNQVSLTCLVKGFYPSDIAVE
WESNGQPENNYDTTPPV LDDSDGSFFLYSDLTVDKSRWQQGNV FSCSV MHE
ALHNHYTQKSLSLSPGGDHCP LGPGRCCRLHTVRASLEDLGWADWVLSR
EVQVTMCIGACPSQFRAANMHAQIKTSLHRLKPDTPAPCCVPASYNPMV
LIQKTD TGVS LQTYDDL LAKDCHCI (SEQ ID NO:276),

- 30 which is encoded by the nucleic acid sequence:

gcacctgaactcctggggggaccgtcagtccttcctcttcccc
caaaacccaaggacaccctcatgatctcccgaccctgaggtcacatgc
gtggtggtggacgtgagccacgaagaccctgaggtcaagttcaactggtg

cgtggacggcgtggaggtgcataattgcaagacaaagccgcgggaggagc
 agtacggcagcacgtaccgtgtgggtcagcgtctgcaccgtcctgcaccag
 gactggctgaatggcaaggagtacaagtgaaggtctccaacaaagccct
 cccagcccccatcgagaaaaccatctccaaagccaaagggcagccccgag
 5 aaccacaggtgtgcaccctgcccccatcccgggaggagatgaccaagaac
 caggtcagcctgacctgctgggtcaaaggcttctatcccagcgacatcgc
 cgtggagtgggagagcaatgggcagccggagaacaactacgacaccacgc
 ctcccgctgctggactccgacggctccttcttctctatagcgacctcacc
 gtggacaagagcaggtggcagcaggggaacgtcttctcatgctccgtgat
 10 gcatgaggctctgcacaaccactacacgcagaagagcctctccctgtctc
 cgggtggagaccactgtccgctcgggccccggcgcttgctgccgtctgcac
 acggtccgcgcgtcgtggaagacctgggctgggccgattgggtgctgtc
 gccacgggaggtgcaagtgaccatgtgcatcggcgcgtgcccgagccagt
 tccggggcggaacatgcacgcgcagatcaagacgagcctgcaccgcctg
 15 aagcccgacacgggtgccagcgccctgctgctgcccggccagctacaatcc
 catgggtgctcattcaaaagaccgacaccggggtgtcgtccagacctatg
 atgacttggttagccaaagactgccactgcata (SEQ ID NO:277) .

In a preferred embodiment, the second polypeptide chain comprises the amino acid sequence:

20 APELLGGPSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVD
 GVEVHNCKTKPREEQYGSTYRVVSVCTV LHQDWLNGKEYKCKVSNKALPA
 PIEKTISKAKGQPREPQVYTLPPCRKEMTKNQVSLTCLVKGFYPSDIAVE
 WESNGQPENNYKTTTPVLKSDGSFFLYSKLTVDKSRWQQGNVFSQVMHE
 ALHNHYTQKSLSLSPGK (SEQ ID NO:274) ,

25 which is encoded by the nucleic acid sequence:

gcacctgaactcctggggggaccgtcagtccttcttcttcccc
 caaaacccaaggacaccctcatgatctcccgaccctgaggtcacatgc
 gtggtggtggacgtgagccacgaagacctgaggtcaagttcaactggta
 cgtggacggcgtggaggtgcataattgcaagacaaagccgcgggaggagc
 30 agtacggcagcacgtaccgtgtgggtcagcgtctgcaccgtgctccaccag
 gactggcttaatgggaaggaatacaagtgttaaggtgtccaacaagccct
 ccccgctcccatcgaaaagaccatctcaaaggcaaaggggaaccaaggg
 aacctcaagtgtacacctgcctccgtgcaggaaggagatgaccaagaac
 caggtcagcctgacttgtctcgtgaagggttctatcccagcgatattgc
 35 tgtggaatgggagtc aaatggccagcccagagaataactacaaaactacc

cacccgtgctgaaatctgatgggtccttcttcttactccaagctgacc
 gtggacaagagccgctggcaacaaggcaatgtcttttagctgctcagtgat
 gcatgagggtctccataatcactacactcagaagtcactgtccctgtctc
 cgggtaaa (SEQ ID NO:278).

- 5 As discussed above, a tetramer is provided comprising two polypeptide chains comprising the sequence of SEQ ID NO:276 and two polypeptide chains comprising the sequence of SEQ ID NO:274.

10 *II.H.11 DhCpmFc(-)(N297G)(A287C)(Y349C)-GDF15(N3D):
 DhCpmFc(+)(N297G)(L306C)(S354C)*

- The designation “DhCpmFc(-)(N297G)(A287C)(Y349C)-GDF15(N3D):DhCpmFc(+)(N297G)(L306C)(S354C)” in the instant disclosure refers to a heterodimer comprising (i) a first polypeptide chain comprising a GDF15(N3D) polypeptide,
 15 the N-terminus of which is linked directly to the C-terminus of a DhCpmFc(-)(N297G)(A287C)(Y349C) domain, and (ii) a second polypeptide chain comprising a DhCpmFc(+)(N297G)(L306C)(S354C) domain.

- In certain embodiments, a tetramer is provided, comprising a dimer of two DhCpmFc(-)(N297G)(A287C)(Y349C)-
 20 GDF15(N3D):DhCpmFc(+)(N297G)(L306C)(S354C) heterodimers in which the two first polypeptide chains of each heterodimer are linked via an interchain disulfide bond between their respective GDF15 regions.

More particularly, in a specific embodiment, the tetramer comprises:

- (a) two DhCpmFc(+)(N297G)(L306C)(Y349C) domains (one each heterodimer)
 25 comprising the sequence of SEQ ID NO:291,
 (b) two DhCpmFc(-)(N297G)(A287C)(S354C) domains (one each heterodimer) comprising the sequence of SEQ ID NO:275,
 and
 (c) two GDF15(N3D) polypeptides (one each heterodimer) comprising the sequence
 30 of SEQ ID NO:52.

In a preferred embodiment, the first polypeptide chain comprises the amino acid sequence:

APELLGGPSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVD
 GVEVHNCKTKPREEQYGSTYRVVSVCTV L H Q D W L N G K E Y K C K V S N K A L P A
 35 P I E K T I S K A K G Q P R E P Q V C T L P P S R E E M T K N Q V S L T C L V K G F Y P S D I A V E
 W E S N G Q P E N N Y D T T P P V L D S D G S F F L Y S D L T V D K S R W Q Q G N V F S C S V M H E
 A L H N H Y T Q K S L S L S P G A R D G D H C P L G P G R C C R L H T V R A S L E D L G W A D W V L

SPREVQVTMCIGACPSQFRAANMHAQIKTSLHRLKPDTPAPCCVPASYN
PMVLIQKTDGTGVSLQTYDDLAKDCHCI (SEQ ID NO:279),

which is encoded by the nucleic acid sequence:

gcacctgaactcctggggggaccgtcagtccttctcttcccc
5 caaaacccaaggacacccctcatgatctcccgacccctgaggtcacatgc
gtggtggtggacgtgagccacgaagaccctgaggtcaagttcaactggta
cgtggacggcgtggaggtgcataattgcaagacaaagccgaggagc
agtacggcagcacgtaccgtgtggtcagcgtctgcaccgtcctgcaccag
gactggctgaatggcaaggagtacaagtgaaggtctccaacaaagccct
10 cccagcccccatcgagaaaaccatctccaaagccaaagggcagccccgag
aaccacaggtgtgcaccctgcccccatcccgaggagatgaccaagaac
caggtcagcctgacctgctgggtcaaaggcttctatcccagcgacatcgc
cgtggagtgggagagcaatgggcagccggagaacaactacgacaccacgc
ctcccgctgctggactccgacggctccttcttctctatagcgacctcacc
15 gtggacaagagcaggtggcagcaggggaacgtcttctcatgctccgtgat
gcatgaggtctgcacaaccactacacgcagaagagcctctccctgtctc
cgggtgcgcgcgacggagaccactgtccgctcgggcccgggcggtgctgc
cgtctgcacacggtccgcgcgtcgtggaagacctgggctgggcccattg
gggtgctgtcggccagggaggtgcaagtgacctgtgcatcggcgcgtgcc
20 cgagccagttccgggcccgaacatgcacgcgcagatcaagacgagcctg
caccgcctgaagcccgcacacgggtgccagcgcctgctgcgtgcccgcag
ctacaatcccatggtgctcattcaaaagaccgacacggggtgtcgtctcc
agacctatgatgacttgtagccaaagactgccactgcata (SEQ ID NO:280).

In a preferred embodiment, the second polypeptide chain comprises the amino acid
25 sequence of SEQ ID NO:274, which is encoded by the nucleic acid sequence of SEQ ID
NO:278.

As discussed above, a tetramer is provided comprising two polypeptide chains
comprising the sequence of SEQ ID NO:279 and two polypeptide chains comprising the
sequence of SEQ ID NO:274.

30

II.H.12 Dh2CpmFc(-)-GDF15(Ndel3):Dh2CpmFc(+)

The designation “Dh2CpmFc(-)-GDF15(Ndel3):Dh2CpmFc(+)” in the instant
disclosure refers to a heterodimer comprising (i) a first polypeptide chain comprising a
GDF15(Ndel3) polypeptide, the N-terminus of which is linked directly to the C-terminus of a
35 Dh2CpmFc(-) domain, and (ii) a second polypeptide chain comprising a Dh2CpmFc(+)
domain.

In certain embodiments, a tetramer is provided, comprising a dimer of two Dh2CpmFc(-)-GDF15(Ndel3):Dh2CpmFc(+) heterodimers in which the two first polypeptide chains of each heterodimer are linked via an interchain disulfide bond between their respective GDF15 regions.

5 More particularly, in a specific embodiment, the tetramer comprises:

(a) two Dh2CpmFc(+) domains (one each heterodimer) comprising the sequence:

PSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVDGVEVHNA
KTKPREEQYNSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPAPIEKTIS
KAKGQPREPQVYTLPPSRKEMTKNQVSLTCLVKGFYPSDIAVEWESNGQP
10 ENNYKTTTPVLKSDGSFFLYSKLTVDKSRWQQGNVFSCSVMHEALHNHYT
QKSLSLSPG (SEQ ID NO:292) ,

(b) two Dh2CpmFc(-) domains (one each heterodimer) comprising the sequence:

PSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVDGVEVHNA
KTKPREEQYNSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPAPIEKTIS
15 KAKGQPREPQVYTLPPSREEMTKNQVSLTCLVKGFYPSDIAVEWESNGQP
ENNYDTTPPVLSDGSFFLYSDLTVDKSRWQQGNVFSCSVMHEALHNHYT
QKSLSLSPG (SEQ ID NO:155) ,

and

(c) two GDF15(Ndel3) polypeptides (one each heterodimer) comprising the sequence
20 of SEQ ID NO:55.

In a preferred embodiment, the first polypeptide chain comprises the amino acid sequence:

PSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVDGVEVHNA
KTKPREEQYNSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPAPIEKTIS
25 KAKGQPREPQVYTLPPSREEMTKNQVSLTCLVKGFYPSDIAVEWESNGQP
ENNYDTTPPVLSDGSFFLYSDLTVDKSRWQQGNVFSCSVMHEALHNHYT
QKSLSLSPGGDHCPLGPGRCCRLHTVRASLEDLGWADWVLSPREVQVTMC
IGACPSQFRAANMHAQIKTSLHRLKPDTPAPCCVPASYNPMVLIQKTD
GVSLQTYDDLLAKDCHCI (SEQ ID NO:157) ,

30 which is encoded by the nucleic acid sequence:

ccgtcagtccttcctcttccccccaaaaccaaggacaccctcatgatctc
ccggacccttgaggtcacatgcgtggtggtggacgtgagccacgaagacc
ctgaggtcaagttcaactggtacgtggacggcgtggaggtgcataatgcc

aagacaaagccgcgggaggagcagtacaacagcacgtaccgtgtggtcag
 cgtcctcaccgtcctgcaccaggactggctgaatggcaaggagtacaagt
 gcaaggttctccaacaaagccctcccagccccatcgagaaaaccatctcc
 aaagccaaagggcagccccgagaaccacaggtgtacaccctgccccatc
 5 cccgggaggagatgaccaagaaccaggtcagcctgacctgacctggtcaaag
 gcttctatcccagcgacatcgccgtggagtgggagagcaatgggcagccg
 gagaacaactacgacaccacgcctcccgtgctggactccgacggctcctt
 ctctctatagcgacctcaccgtggacaagagcaggtggcagcagggga
 acgtcttctcatgctccgtgatgcatgaggctctgcacaaccactacacg
 10 cagaagagcctctccctgtctccgggtggagaccactgtccgctcggggc
 cgggcgttgctgccgtctgcacacgggtccgcgcgtcgtggaagacctgg
 gctggggccgattgggtgctgtcgccacgggaggtgcaagtgacctgtgc
 atcgggcgcgtgcccagccagttccgggcggcaaacatgcacgcgcagat
 caagacgagcctgcaccgcctgaagcccagacacggtgccagcgcctgct
 15 gcgtgcccgcagctacaatcccatggtgctcattcaaaagaccgacacc
 ggggtgtcgtccagacctatgatgacttggttagccaaagactgccactg
 cata (SEQ ID NO:156).

In a preferred embodiment, the second polypeptide chain comprises the amino acid sequence:

20 PSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVDGVEVHNA
 KTKPREEQYNSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPAPIEKTIS
 KAKGQPREPQVYTLPPSRKEMTKNQVSLTCLVKGFYPSDIAVEWESNGQP
 ENNYKTTTPVLKSDGSFFLYSKLTVDKSRWQQGNVFSVSMHEALHNHYT
 QKSLSLSPGK (SEQ ID NO:154),

25 which is encoded by the nucleic acid sequence:

ccgtcagtccttctcttccccccaaaacccaaggacaccctcatgatctc
 ccggacccttgaggtcacatgcgtggtggtggactgagccacgaagacc
 ctgaggtcaagttcaactggtacgtggacggcgtggaggtgcataatgcc
 aagacaaagccgcgggaggagcagtacaacagcacgtaccgtgtggtcag
 30 cgtcctcaccgtcctgcaccaggactggctgaatggcaaggagtacaagt
 gcaaggttctccaacaaagccctcccagccccatcgagaaaaccatctcc
 aaagccaaagggcagccccgagaaccacaggtgtacaccctgccccatc
 ccggaaggagatgaccaagaaccaggtcagcctgacctgacctggtcaaag
 gcttctatcccagcgacatcgccgtggagtgggagagcaatgggcagccg
 35 gagaacaactacaagaccacgcctcccgtgctgaagtccgacggctcctt

cttcctctatagcaagctcaccgtggacaagagcaggtggcagcagggga
acgtctttctcatgctccgtgatgcatgaggtcttgacacaccactacacg
cagaagagcctctccctgtctccgggtaaatga (SEQ ID NO:158)

As discussed above, a tetramer is provided comprising two polypeptide chains
5 comprising the sequence of SEQ ID NO:157 and two polypeptide chains comprising the
sequence of SEQ ID NO:154.

II.H.13 Dh2CpmFc(-)-GDF15(N3D):Dh2CpmFc(+)

The designation “Dh2CpmFc(-)-GDF15(N3D):Dh2CpmFc(+)” in the instant
10 disclosure refers to a heterodimer comprising (i) a first polypeptide chain comprising a
GDF15(N3D) polypeptide, the N-terminus of which is linked directly to the C-terminus of a
Dh2CpmFc(-) domain by a peptide bond, and (ii) a second polypeptide chain comprising a
Dh2CpmFc(+) domain.

In certain embodiments, a tetramer is provided, comprising a dimer of two
15 Dh2CpmFc(-)-GDF15(N3D):DhCpmFc(+) heterodimers in which the two first polypeptide
chains of each heterodimer are linked via an interchain disulfide bond between their
respective GDF15 regions.

More particularly, in a specific embodiment, the tetramer comprises:

(a) two Dh2CpmFc(+) domains (one each heterodimer) comprising the sequence of
20 SEQ ID NO:292,

(b) two Dh2CpmFc(-) domains (one each heterodimer) comprising the sequence of
SEQ ID NO:155, and

(c) two GDF15(N3D) polypeptides (one each heterodimer) comprising the sequence
of SEQ ID NO:52.

25 In a preferred embodiment, the first polypeptide chain comprises the amino acid
sequence:

PSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVDGVEVHNA
KTKPREEQYNSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPAPIEKTIS
KAKGQPREPQVYTLPPSREEMTKNQVSLTCLVKGFYPSDIAVEWESNGQP
30 ENNYDTTPPVLDSDGSFFLYSDLTVDKSRWQQGNVFSVMHEALHNHYT
QKSLSLSPGARDGDHCPLGPRCCRLHTVRASLEDLGWADWVLSPREVQV
TMCIGACPSQFRAANMHAQIKTSLHRLKPDTPAPCCVPASYNPMVLIQK
TDTGVSLQTYDDLAKDCHCI (SEQ ID NO:160),

which is encoded by the nucleic acid sequence:

ccgtcagtccttctcttccccccaaaacccaaggacaccctcatgatctc
 ccggacccctgaggtcacatgcgtggtggtggacgtgagccacgaagacc
 ctgaggtcaagttcaactggtacgtggacggcggtggaggtgcataatgcc
 aagacaaagccgcgggaggagcagtacaacagcacgtaccgtgtggtcag
 5 cgtcctcaccgtcctgcaccaggactggctgaatggcaaggagtacaagt
 gcaagggtctccaacaaagccctcccagccccatcgagaaaaccatctcc
 aaagccaaagggcagccccgagaaccacaggtgtacaccctgccccatc
 ccgggaggagatgaccaagaaccaggtcagcctgacctgctggtcaaag
 gcttctatcccagcgacatcgccgtggagtgaggagcaatgggcagccg
 10 gagaacaactacgacaccacgcctcccgctgctggactccgacggctcctt
 ctctctctatagcgacctcaccgtggacaagagcaggtggcagcagggga
 acgtctcttcatgctccgtgatgcatgaggtctgcacaaccactacacg
 cagaagagcctctccctgtctccgggtgcgcgcgacggagaccactgtcc
 gctcgggccccggcggttgctgccgtctgcacacggctccgcgcgtcgtg
 15 aagacctgggctgggcccattgggtgctgtcgccacgggaggtgcaagtg
 accatgtgcatcggcgcgtgcccagccagttccgggaggcaaacatgca
 cgcgcagatcaagacgagcctgcaccgcctgaagcccgacacgggtgccag
 cgcctgctgctgcccgcagctacaatcccatggtgctcattcaaaag
 accgacaccggggtgtcgtccagacctatgatgacttgtagccaaaga
 20 ctgccactgcata (SEQ ID NO:159).

In a preferred embodiment, the second polypeptide chain comprises the amino acid sequence of SEQ ID NO:154, which is encoded by the nucleic acid sequence of SEQ ID NO:158.

As discussed above, a tetramer is provided comprising two polypeptide chains
 25 comprising the sequence of SEQ ID NO:160 and two polypeptide chains comprising the
 sequence of SEQ ID NO:154.

II.H.14 Dh2CpmFc(-)(Y349C)-GDF15(Ndel3):Dh2CpmFc(+)(S354C)

The designation “Dh2CpmFc(-)(Y349C)-GDF15(Ndel3):Dh2CpmFc(+)(S354C)” in
 30 the instant disclosure refers to a heterodimer comprising (i) a first polypeptide chain
 comprising a GDF15(Ndel3) polypeptide, the N-terminus of which is linked directly to the C-
 terminus of a Dh2CpmFc(-)(Y349C) domain, and (ii) a second polypeptide chain comprising
 a Dh2CpmFc(+)(S354C) domain. The cysteine clamp mutations allow the first and second
 polypeptide chains to be linked via an interchain disulfide bond between C349 of the first
 35 polypeptide chain and C354 of the second polypeptide chain.

In certain embodiments, a tetramer is provided, comprising a dimer of two Dh2CpmFc(-)(Y349C)-GDF15(Nde13):Dh2CpmFc(+)(S354C) heterodimers in which the two first polypeptide chains of each heterodimer are linked via an interchain disulfide bond between their respective GDF15 regions.

5 More particularly, in a specific embodiment, the tetramer comprises:

(a) two Dh2CpmFc(+)(S354C) domains (one each heterodimer) comprising the sequence:

PSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVDGVEVHNA
KTKPREEQYNSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPAPIEKTIS
10 KAKGQPREPQVYITLPPCRKEMTKNQVSLTCLVKGFYPSDIAVEWESNGQP
ENNYKTTTPVLKSDGSFFLYSKLTVDKSRWQQGNVFSVSMHEALHNHYT
QKSLSLSPG (SEQ ID NO:293),

(b) two Dh2CpmFc(-)(Y349C) domains (one each heterodimer) comprising the sequence:

15 PSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVDGVEVHNA
KTKPREEQYNSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPAPIEKTIS
KAKGQPREPQVCTLPSPREEMTKNQVSLTCLVKGFYPSDIAVEWESNGQP
ENNYDTTPPVLDSGSDGSFFLYSDLTVDKSRWQQGNVFSVSMHEALHNHYT
QKSLSLSPG (SEQ ID NO:162),

20 and

(c) two GDF15(Nde13) polypeptides (one each heterodimer) comprising the sequence of SEQ ID NO:55.

In a preferred embodiment, the first polypeptide chain comprises the amino acid sequence:

25 PSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVDGVEVHNA
KTKPREEQYNSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPAPIEKTIS
KAKGQPREPQVCTLPSPREEMTKNQVSLTCLVKGFYPSDIAVEWESNGQP
ENNYDTTPPVLDSGSDGSFFLYSDLTVDKSRWQQGNVFSVSMHEALHNHYT
QKSLSLSPGSDHCPGRCRLHTVRASLEDLGWADWVLSPREVQVTMC
30 IGACPSQFRAANMHAQIKTSLHRLKPDTPAPCCVPASYNPMVLIQKTD
GVSLQTYDDLLAKDCHCI (SEQ ID NO:164),

which is encoded by the nucleic acid sequence:

ccgtcagtccttctcttccccccaaaacccaaggacaccctcatgatctc
 ccggacccctgaggtcacatgcgtggtggtggacgtgagccacgaagacc
 ctgaggtcaagttcaactggtacgtggacggcgtggaggtgcataatgcc
 aagacaaagccgcgggaggagcagtacaacagcacgtaccgtgtggtcag
 5 cgtcctcacctcctgcaccaggactggctgaatggcaaggagtacaagt
 gcaaggtctccaacaagccctcccagccccatcgagaaaaccatctcc
 aaagccaaagggcagccccgagaaccacaggtgtgcaccctgccccatc
 ccgggaggagatgaccaagaaccaggtcagcctgacctgcctggtcaaag
 gcttctatcccagcgacatcgccgtggagtgaggagcaatgggcagccg
 10 gagaacaactacgacaccacgcctcccgtgctggactccgacggctcctt
 ctctctatagcgacctcaccgtggacaagagcaggtggcagcagggga
 acgtcttctcatgctccgtgatgcatgaggtctgcacaaccactacacg
 cagaagagcctctccctgtctccgggtggagaccactgtccgctcgggcc
 cgggcgttgctgccgtctgcacacgggtccgcgcgtcgctggaagacctgg
 15 gctggggccgattgggtgctgtcgccacgggaggtgcaagtgacctgtgc
 atcgggcgcgtgcccagccagttccggggcggcaaacatgcacgcgcagat
 caagacgagcctgcaccgcctgaagcccgacacgggtgccagcgcctgtc
 gcggtgcccgccagctacaatcccattggtgctcattcaaaagaccgacacc
 ggggtgtcgtccagacctatgatgacttgtagccaaagactgccactg
 20 cata (SEQ ID NO:163).

In a preferred embodiment, the second polypeptide chain comprises the amino acid sequence:

PSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVDGVEVHNA
 KTKPREEQYNSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPAPIEKTIS
 25 KAKGQPREPQVYTLPPCRKEMTKNQVSLTCLVKGFYPSDIAVEWESNGQP
 ENNYKTTTPVLKSDGSFFLYSKLTVDKSRWQQGNVFSVSMHEALHNHYT
 QKSLSLSPGK (SEQ ID NO:161),

which is encoded by the nucleic acid sequence:

ccatccgtgttctctgtttctccaaagccgaaggacaccctgatgatctc
 30 aagaactccggaagtgacttgcgtcgctcgaggacgtgtcacatgaggatc
 cagaggtcaagttcaattggtatgtggacggagtggaagtgcataacgcc
 aagaccaaaccgccgaagaacagtacaatagcacctaccgcgtggtgag
 cgtccttactgtgctccaccaggactggcttaatgggaaggaataacaagt
 gtaaggtgtccaacaaggccctcccgcctcccatcgaaaagaccatctca
 35 aaggcaaaaggggcaaccaagggaacctcaagtgtacaccctgcctccgtg

caggaaggagatgaccaagaaccagggtcagcctgacttgtctcgtgaagg
 gcttctatcccagcgatattgctgtggaatgggagtcaaattggccagccc
 gagaataactacaaaactacccccaccgtgctgaaatctgatgggtcctt
 cttccttttactccaagctgaccgtggacaagagccgctggcaacaaggca
 5 atgtcttttagctgctcagtgatgcatgaggtctccataatcactacact
 cagaagtcactgtccctgtctccgggtaaa (SEQ ID NO:165) .

As discussed above, a tetramer is provided comprising two polypeptide chains comprising the sequence of SEQ ID NO:164 and two polypeptide chains comprising the sequence of SEQ ID NO:161.

10

II.H.15 Dh2CpmFc(-)(Y349C)-GDF15(N3D):Dh2CpmFc(+)(S354C)

The designation “Dh2CpmFc(-)(Y349C)-GDF15(N3D):Dh2CpmFc(+)(S354C)” in the instant disclosure refers to a heterodimer comprising (i) a first polypeptide chain comprising a GDF15(N3D) polypeptide, the N-terminus of which is linked directly to the C-terminus of a Dh2CpmFc(-)(Y349C) domain, and (ii) a second polypeptide chain comprising a Dh2CpmFc(+)(S354C) domain. The cysteine clamp mutations allow the first and second polypeptide chains to be linked via an interchain disulfide bond between C349 of the first polypeptide chain and C354 of the second polypeptide chain.

In certain embodiments, a tetramer is provided, comprising a dimer of two Dh2CpmFc(-)(Y349C)-GDF15(N3D):Dh2CpmFc(+)(S354C) heterodimers in which two first polypeptide chains of each heterodimer are linked via an interchain disulfide bond between their respective GDF15 regions.

More particularly, in a specific embodiment, the tetramer comprises:

(a) two Dh2CpmFc(+)(S354C) domains (one each heterodimer) comprising the sequence of SEQ ID NO:293,

(b) two Dh2CpmFc(-)(Y349C) domains (one each heterodimer) comprising the sequence of SEQ ID NO:162, and

(c) two GDF15(N3D) polypeptides (one each heterodimer) comprising the sequence of SEQ ID NO:52.

In a preferred embodiment, the first polypeptide chain comprises the amino acid sequence:

PSVFLFPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVDGVEVHNA
 KTKPREEQYNSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPAPIEKTIS
 KAKGQPREPQVCTLPSPREEMTKNQVSLTCLVKGFYPSDIAVEWESNGQP
 35 ENNYDTTPPVLDSDGSFFLYSDLTVDKSRWQQGNVFSQVMHEALHNHYT
 QKSLSLSPGARDGDHCPLGPGRCCLHTVRSLEDLGWADWVLSPREVQV

TMCIGACPSQFRAANMHAQIKTSLHRLKPDTPAPCCVPASYNPMVLIQK
TDTGVSLQTYDDLLAKDCHCI (SEQ ID NO:167),

which is encoded by the nucleic acid sequence:

ccgtcagtccttctcttccccccaaaacccaaggacaccctcatgatctc
5 cccgacccctgaggtcacatgcgtggtggtggacgtgagccacgaagacc
ctgaggtcaagttcaactggtacgtggacggcgtggaggtgcataatgcc
aagacaaagccgcgggaggagcagtacaacagcacgtaccgtgtggtcag
cgtcctcaccgtcctgcaccaggactggctgaatggcaaggagtacaagt
gcaaggtctccaacaaagccctcccagccccatcgagaaaaccatctcc
10 aaagccaaagggcagccccgagaaccacaggtgtgcaccctgccccatc
ccgggaggagatgaccaagaaccaggtcagcctgacctgcctggtcaaag
gcttctatcccagcgacatcgccgtggagtgaggagcaatgggcagccg
gagaacaactacgacaccacgcctcccgtgctggactccgacggctcctt
cttctctatagcgacctcaccgtggacaagagcaggtggcagcagggga
15 acgtcttctcatgctccgtgatgcatgaggtctgcacaaccactacacg
cagaagagcctctccctgtctccgggtgcgcgcgacggagaccactgtcc
gctcgggccccggcggttgcctgctgcacacggctccgcgcgtcgctgg
aagacctgggctgggcccattgggtgctgctgccacgggaggtgcaagtg
accatgtgcatcggcgcgtgcccagccagttccgggcccgaacatgca
20 cgcgcagatcaagacgagcctgcaccgcctgaagcccgcacgggtgccag
cgccctgctgctgcccgccagctacaatcccatggtgctcattcaaaag
accgacaccggggtgctcgtccagacctatgatgacttgtagccaaaga
ctgccactgcata (SEQ ID NO:166).

In a preferred embodiment, the second monomer comprises the amino acid sequence
25 of SEQ ID NO:161, which is encoded by the nucleic acid sequence of SEQ ID NO:165.

As discussed above, a tetramer is provided comprising two polypeptide chains
comprising the sequence of SEQ ID NO:167 and two polypeptide chains comprising the
sequence of SEQ ID NO:161.

30 II.H.16 *CpmFc(-)(N297G)-GDF15(Ndel3):CpmFc(+)(N297G)*

The designation “CpmFc(-)(N297G)-GDF15(Ndel3):DhCpmFc(+)(N297G)” in the
instant disclosure refers to a heterodimer comprising (i) a first polypeptide chain comprising a
GDF15(Ndel3) polypeptide, the N-terminus of which is linked directly to the C-terminus of a
CpmFc(-)(N297G) domain, and (ii) a second polypeptide chain comprising a
35 CpmFc(+)(N297G) domain.

In certain embodiments, a tetramer is provided, comprising a dimer of two CpmFc(-)(N297G)-GDF15(Nde13):CpmFc(+)(N297G) heterodimers in which the two first polypeptide chains of each heterodimer are linked via an interchain disulfide bond between their respective GDF15 regions.

5 More particularly, in a specific embodiment, the tetramer comprises:

(a) two CpmFc(+)(N297G) domains (one each heterodimer) comprising the sequence (part of the hinge region in parentheses):

(DKTHTCPPCP) APELLGGPSVFLFPPKPKDTLMISRTPEVTCVVVDVSH
EDPEVKFNWYVDGVEVHNAKTKPREEQYGSTYRVVSVLTVLHQDWLNGKE
10 YKCKVSNKALPAPIEKTISKAKGQPREPQVYTLPPSRKEMTKNQVSLTCL
VKGFIYPSDIAVEWESNGQPENNYKTTPPVLSKSDGSFFLYSKLTVDKSRWQ
QGNVFSCSVMHEALHNHYTQKSLSLSPG (SEQ ID NO:294),

(b) two CpmFc(-)(N297G) domains (one each heterodimer) comprising the sequence (part of the hinge region in parentheses):

15 (DKTHTCPPCP) APELLGGPSVFLFPPKPKDTLMISRTPEVTCVVVDVSH
EDPEVKFNWYVDGVEVHNAKTKPREEQYGSTYRVVSVLTVLHQDWLNGKE
YKCKVSNKALPAPIEKTISKAKGQPREPQVYTLPPSREEMTKNQVSLTCL
VKGFIYPSDIAVEWESNGQPENNYDTTPPVLDSDGSFFLYSDLTVDKSRWQ
QGNVFSCSVMHEALHNHYTQKSLSLSPG (SEQ ID NO:169),

20 and

(c) two GDF15(Nde13) polypeptides (one each heterodimer) comprising the sequence of SEQ ID NO:55.

In a preferred embodiment, the first polypeptide chain comprises the amino acid sequence:

25 (DKTHTCPPCP) APELLGGPSVFLFPPKPKDTLMISRTPEVTCVVVDVSH
EDPEVKFNWYVDGVEVHNAKTKPREEQYGSTYRVVSVLTVLHQDWLNGKE
YKCKVSNKALPAPIEKTISKAKGQPREPQVYTLPPSREEMTKNQVSLTCL
VKGFIYPSDIAVEWESNGQPENNYDTTPPVLDSDGSFFLYSDLTVDKSRWQ
QGNVFSCSVMHEALHNHYTQKSLSLSPGGDHCPGPRCCRLHTVRSLE
30 DLGWADWVLSPREVQVTMCIGACPSQFRAANMHAQIKTSLHRLKPDTPVA
PCCVPASYNPMVLIQKTDGTGVSQTYYDILLAKDCHCI (SEQ ID NO:171),

which is encoded by the nucleic acid sequence:

gacaaaactcacacatgcccaccgtgcccagcacctgaactcc
 tgggggggaccgtcagtccttctcttccccccaaaacccaaggacaccctc
 atgatctcccggaccctgaggtcacatgcgtggtggtggacgtgagcca
 cgaagaccctgaggtcaagttcaactggtacgtggacggcgtggaggtgc
 5 ataatgccaagacaaagccgcgggaggagcagtacggcagcacgtaccgt
 gtggtcagcgtcctcaccgtcctgcaccaggactggctgaatggcaagga
 gtacaagtgcagggtctccaacaaagccctcccagcccccatcgagaaaa
 ccatctccaaagccaaagggcagccccgagaaccacaggtgtacaccctg
 ccccatcccgggaggagatgaccaagaaccaggtcagcctgacctgcct
 10 ggtcaaaggcttctatcccagcgacatcgccgtggagtgggagagcaatg
 ggcagccggagaacaactacgacaccacgcctcccgtgctggactccgac
 ggctccttcttctctatagcgacctcaccgtggacaagagcaggtggca
 gcaggggaacgtcttctcatgctccgtgatgcatgaggctctgcacaacc
 actacacgcagaagagcctctccctgtctccgggtggagaccactgtccg
 15 ctccggggcccggttgctgccgtctgcacacgggtccgcgcgtcgctgga
 agacctgggctgggcccattgggtgctgtcgccacgggaggtgcaagtga
 ccatgtgcatcggcgcgtgcccagaccagttccggggcggaacatgcac
 gcgcagatcaagacgagcctgcaccgcctgaagcccgcacacgggtgccagc
 gccctgctgctgcccgcagctacaatcccatggtgctcattcaaaaga
 20 ccgacaccggggtgctgctccagacctatgatgacttgtagccaaagac
 tgccactgcata (SEQ ID NO:170).

In a preferred embodiment, the second polypeptide chain comprises the amino acid sequence:

(DKHTCPCP) APELLGGPSVFLFPPKPKDTLMISRTPEVTCVVVDVSH
 25 EDPEVKFNWYVDGVEVHNAKTKPREEQYGSTYRVVSVLTVLHQDWLNGKE
 YKCKVSNKALPAPIEKTISKAKGQPREPQVYTLPPSRKEMTKNQVSLTCL
 VKGFYPSDIAVEWESNGQPENNYKTTPPVLKSDGSFFLYSKLTVDKSRWQ
 QGNVFSCSVMHEALHNHYTQKSLSLSPGK (SEQ ID NO:168),

which is encoded by the nucleic acid sequence:

30 gacaaaactcacacatgcccaccgtgcccagcacctgaactcc
 tgggggggaccgtcagtccttctcttccccccaaaacccaaggacaccctc
 atgatctcccggaccctgaggtcacatgcgtggtggtggacgtgagcca
 cgaagaccctgaggtcaagttcaactggtacgtggacggcgtggaggtgc
 ataatgccaagacaaagccgcgggaggagcagtacggcagcacgtaccgt
 35 gtggtcagcgtcctcaccgtcctgcaccaggactggctgaatggcaagga

gtacaagtgcaagggtctccaacaaagccctcccagcccccatcgagaaaa
 ccatctccaaagccaaagggcagccccgagaaccacaggtgtacacctg
 cccccatcccgaaggagatgaccaagaaccaggtcagcctgacctgcct
 ggtcaaaggcttctatcccagcgacatcgccgtggagtgggagagcaatg
 5 ggcagccggagaacaactacaagaccacgcctcccgtgctgaagtccgac
 ggctccttcttctctatagcaagctcaccgtggacaagagcaggtggca
 gcaggggaacgtcttctcatgctccgtgatgcatgaggctctgcacaacc
 actacacgcagaagagcctctccctgtctccgggtaaa (SEQ ID NO:172) .

As discussed above, a tetramer is provided comprising two polypeptide chains
 10 comprising the sequence of SEQ ID NO:171 and two polypeptide chains comprising the
 sequence of SEQ ID NO:168.

II.H.17 CpmFc(-)(N297G)-GDF15(N3D):CpmFc(+)(N297G)

The designation “CpmFc(-)(N297G)-GDF15(N3D):DhCpmFc(+)(N297G)” in the
 15 instant disclosure refers to a heterodimer comprising (i) a first polypeptide chain comprising a
 GDF15(N3D) polypeptide, the N-terminus of which is linked directly to the C-terminus of a
 CpmFc(-)(N297G) domain, and (ii) a second polypeptide chain comprising a
 CpmFc(+)(N297G) domain.

In certain embodiments, a tetramer is provided, comprising a dimer of two
 20 CpmFc(-)(N297G)-GDF15(N3D):CpmFc(+)(N297G) heterodimers in which the two first
 polypeptide chains of each heterodimer are linked via an interchain disulfide bond between
 their respective GDF15 regions.

More particularly, in a specific embodiment, the tetramer comprises:

(a) two CpmFc(+)(N297G) domains (one each heterodimer) comprising the sequence
 25 of SEQ ID NO:294,

(b) two CpmFc(-)(N297G) domains (one each heterodimer) comprising the sequence
 of SEQ ID NO:169, and

(c) two GDF15(N3D) polypeptides (one each heterodimer) comprising the sequence
 of SEQ ID NO:52.

30 In a preferred embodiment, the first polypeptide chain comprises the amino acid
 sequence:

(DKTHTCPPCP) APELLGGPSVFLFPPKPKDTLMISRTPEVTCVVVDVSH
 EDPEVKFNWYVDGVEVHNAKTKPREEQYGSTYRVVSVLTVLHQDWLNGKE
 YKCKVSNKALPAPIEKTISKAKGQPREPQVYTLPPSREEMTKNQVSLTCL
 35 VKGFYPSDIAVEWESNGQPENNYDTTPPVLDSDGSFFLYSDLTVDKSRWQ
 QGNVFSCSVMHEALHNHYTQKSLSLSPGARDGDHCP LGPGRCCRLHTVRA

SLEDLGWADWVLSPREVQVTMCIGACPSQFRAANMHAQIKTSLHRLKPDT
VPAPCCVPASYNPMVLIQKTDGTGVSLLQTYDDLLAKDCHCI (SEQ ID NO:174),

which is encoded by the nucleic acid sequence:

gacaaaactcacacatgcccaccgtgcccagcacctgaactcc
5 tgggggggaccgtcagtccttctcttccccccaaaacccaaggacaccctc
atgatctcccgaccctgaggtcacatgcgtggtggtggacgtgagcca
cgaagaccctgaggtcaagttcaactggtacgtggacggcgtggaggtgc
ataatgccaagacaaagccgcgggaggagcagtacggcagcacgtaccgt
gtggtcagcgtcctcaccgtcctgcaccaggactggctgaatggcaagga
10 gtacaagtgcagggtctccaacaaagccctcccagcccccatcgagaaaa
ccatctccaaagccaaagggcagccccgagaaccacaggtgtacaccctg
cccccatcccgggaggagatgaccaagaaccaggtcagcctgacctgcct
ggtcaaaggcttctatcccagcgacatcgccgtggagtgggagagcaatg
ggcagccggagaacaactacgacaccacgcctcccgtgctggactccgac
15 ggctccttcttctctatagcgacctcaccgtggacaagagcaggtggca
gcaggggaacgtcttctcatgctccgtgatgcatgaggctctgcacaacc
actacacgcagaagagcctctccctgtctccgggtgcgcgcgacggagac
cactgtccgctcggggccggcggttctgctgcccgtctgcacacgggtccgcgc
gtcgtggaagacctgggctgggcccattgggtgctgctgccacgggagg
20 tgcaagtgacctgtgcatcggcgcgtgcccagaccagttccgggggga
aacatgcacgcgcagatcaagacgagcctgcaccgctgaagcccgacac
gggtgccagcgccctgctgctgcccgcagctacaatcccatggtgctca
ttcaaaagaccgacaccggggtgctcgtccagacctatgatgacttgta
gccaaagactgccactgcata (SEQ ID NO:173).

25 In a preferred embodiment, the second polypeptide chain comprises the amino acid sequence of SEQ ID NO:168, which is encoded by the nucleic acid sequence comprising the sequence of SEQ ID NO:172.

As discussed above, a tetramer is provided comprising two polypeptide chains comprising the sequence of SEQ ID NO:174 and two polypeptide chains comprising the
30 sequence of SEQ ID NO:168.

II.H.18 Dh2CpmFc(-)(N297G)-GDF15(Ndel3):Dh2CpmFc(+)(N297G)

The designation “Dh2CpmFc(-)(N297G)-GDF15(Ndel3):Dh2CpmFc(+)(N297G)” in the instant disclosure refers to a heterodimer comprising (i) a first polypeptide chain
35 comprising a GDF15(Ndel3) polypeptide, the N-terminus of which is linked directly to the C-

terminus of a Dh2CpmFc(-)(N297G) domain, and (ii) a second polypeptide chain comprising a Dh2CpmFc(+)(N297G) domain.

In certain embodiments, a tetramer is provided, comprising a dimer of two Dh2CpmFc(-)(N297G)-GDF15(Nde13):Dh2CpmFc(+)(N297G) heterodimers in which the two first polypeptide chains of each heterodimer are linked via an interchain disulfide bond between their respective GDF15 regions.

More particularly, in a specific embodiment, the tetramer comprises:

(a) two Dh2CpmFc(+)(N297G) domains (one each heterodimer) comprising the sequence:

10 PSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVDGVEVHNA
KTKPREEQYGSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPAPIEKTIS
KAKGQPREPQVYTLPPSRKEMTKNQVSLTCLVKGFYPSDIAVEWESNGQP
ENNYKTTTPVLKSDGSFFLYSKLTVDKSRWQQGNVFSCSVMHEALHNHYT
QKSLSLSPG (SEQ ID NO:295),

15 (b) two Dh2CpmFc(-)(N297G) domains (one each heterodimer) comprising the sequence:

PSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVDGVEVHNA
KTKPREEQYGSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPAPIEKTIS
KAKGQPREPQVYTLPPSREEMTKNQVSLTCLVKGFYPSDIAVEWESNGQP
20 ENNYDTTPVLDSDGSFFLYSDLTVDKSRWQQGNVFSCSVMHEALHNHYT
QKSLSLSPG (SEQ ID NO:176),

and

(c) two GDF15(Nde13) polypeptides (one each heterodimer) comprising the sequence of SEQ ID NO:55.

25 In a preferred embodiment, the first polypeptide chain comprises the amino acid sequence:

PSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVDGVEVHNA
KTKPREEQYGSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPAPIEKTIS
KAKGQPREPQVYTLPPSREEMTKNQVSLTCLVKGFYPSDIAVEWESNGQP
30 ENNYDTTPVLDSDGSFFLYSDLTVDKSRWQQGNVFSCSVMHEALHNHYT
QKSLSLSPG GDHCPGPRCCRLHTVRASLEDLGWADWVLSPREVQVTMC
IGACPSQFRAANMHAQIKTSLHRLKPDTPAPCCVPASYNPMVLIQKTD
GVSLQTYDDLLAKDCHCI (SEQ ID NO:178),

which is encoded by the nucleic acid sequence:

ccgtcagtccttctcttccccccaaaacccaaggacaccctcatgatctc
 ccggacccctgaggtcacatgcgtggtggtggacgtgagccacgaagacc
 ctgaggtcaagttcaactggtacgtggacggcggtggaggtgcataatgcc
 aagacaaagccgcgggaggagcagtacgggagcacgtaccgtgtggtcag
 5 cgtcctcaccgtcctgcaccaggactggctgaatggcaaggagtacaagt
 gcaaggtctccaacaaagccctcccagcccccatcgagaaaaccatctcc
 aaagccaaagggcagccccgagaaccacaggtgtacaccctgcccccatc
 ccgggaggagatgaccaagaaccaggtcagcctgacctgcctggtcaaag
 gcttctatcccagcgacatcgccgtggagtgaggagcaatgggcagccg
 10 gagaacaactacgacaccacgcctcccgtgctggactccgacggctcctt
 ctctctctatagcgacctcaccgtggacaagagcaggtggcagcagggga
 acgtctctctcatgctccgtgatgcatgaggtctgcacaaccactacacg
 cagaagagcctctccctgtctccgggtggagaccactgtccgctcgggccc
 cgggcggttgctgccgtctgcacacgggtccgcgcgtcgctggaagacctgg
 15 gctggggccgattgggtgctgtcgccacgggaggtgcaagtgacctatgtgc
 atcgggcgcgtgcccagccagttccggggcggcaaacatgcacgcgcagat
 caagacgagcctgcaccgcctgaagcccgacacgggtgccagcgccttget
 gcggtgcccgccagctacaatcccattggtgctcattcaaaagaccgacacc
 ggggtgtcgtccagacctatgatgacttggttagccaaagactgccactg
 20 cata (SEQ ID NO:177) .

In a preferred embodiment, the second polypeptide chain comprises the amino acid sequence:

PSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVDGVEVHNA
 KTKPREEQYGSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPAPIEKTIS
 25 KAKGQPREPQVYTLPPSRKEMTKNQVSLTCLVKGFYPSDIAVEWESNGQP
 ENNYKTTTPVLKSDGSFFLYSKLTVDKSRWQQGNVFSQSVMEALHNHYT
 QKSLSLSPGK (SEQ ID NO:175) ,

which is encoded by the nucleic acid sequence:

ccgtcagtccttctcttccccccaaaacccaaggacaccctcatgatctc
 30 ccggacccctgaggtcacatgcgtggtggtggacgtgagccacgaagacc
 ctgaggtcaagttcaactggtacgtggacggcggtggaggtgcataatgcc
 aagacaaagccgcgggaggagcagtacgggagcacgtaccgtgtggtcag
 cgtcctcaccgtcctgcaccaggactggctgaatggcaaggagtacaagt
 gcaaggtctccaacaaagccctcccagcccccatcgagaaaaccatctcc
 35 aaagccaaagggcagccccgagaaccacaggtgtacaccctgcccccatc

ccggaaggagatgaccaagaaccaggtcagcctgacctgcctgggtcaaag
 gcttctatcccagcgacatcgccgtggagtgggagagcaatgggcagccg
 gagaacaactacaagaccacgcctcccgtgctgaagtccgacggctcctt
 cttcctctatagcaagctcacggtggacaagagcaggtggcagcagggga
 5 acgtcttctcatgctccgtgatgcatgaggtctgcacaaccactacacg
 cagaagagcctctccctgtctccgggtaaatga (SEQ ID NO:179).

As discussed above, a tetramer is provided comprising two polypeptide chains comprising the sequence of SEQ ID NO:178 and two polypeptide chains comprising the sequence of SEQ ID NO:175.

10

II.H.19 Dh2CpmFc(-)(N297G)-GDF15(N3D):Dh2CpmFc(+)(N297G)

The designation “Dh2CpmFc(-)(N297G)-GDF15(N3D):DhCpmFc(+)(N297G)” in the instant disclosure refers to a heterodimer comprising (i) a first polypeptide chain comprising a GDF15(N3D) polypeptide, the N-terminus of which is linked directly to the C-terminus of a Dh2CpmFc(-)(N297G) domain, and (ii) a second polypeptide chain comprising a Dh2CpmFc(+)(N297G) domain.

In certain embodiments, a tetramer is provided, comprising a dimer of two Dh2CpmFc(-)(N297G)-GDF15(N3D):Dh2CpmFc(+)(N297G) heterodimers in which the two first polypeptide chains of each heterodimer are linked via an interchain disulfide bond between their respective GDF15 regions.

More particularly, in a specific embodiment, the tetramer comprises:

- (a) two Dh2CpmFc(+)(N297G) domains (one each heterodimer) comprising the sequence of SEQ ID NO:295,
- (b) two Dh2CpmFc(-)(N297G) domains (one each heterodimer) comprising the sequence of SEQ ID NO:176, and
- (c) two GDF15(N3D) polypeptides (one each heterodimer) comprising the sequence of SEQ ID NO:52.

In a preferred embodiment, the first polypeptide chain comprises the amino acid sequence:

PSVFLFPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVDGVEVHNA
 KTKPREEQYGSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPAPIEKTIS
 KAKGQPREPQVYTLPPSREEMTKNQVSLTCLVKGFYPSDIAVEWESNGQP
 ENNYDTTPPVLDSDGSFFLYSDLTVDKSRWQQGNVFCFSVMHEALHNHYT
 QKSLSLSPGARDGDHCPGPRCCRLHTVRASLEDLGWADWVLSPREVQV
 35 TMCIGACPSQFRAANMHAQIKTSLHRLKPDTPAPCCVPASYNPMVLIQK
 TDTGVSLQTYDDLLAKDCHCI (SEQ ID NO:181),

which is encoded by the nucleic acid sequence:

```

ccgtcagtccttccctcttccccccaaaacccaaggacaccctcatgatctc
ccggacccctgaggtcacatgcgtggtggtggacgtgagccacgaagacc
ctgaggtcaagttcaactggtacgtggacggcgtggaggtgcataatgcc
5 aagacaaagccgcgggaggagcagtagcgggagcacgtaccgtgtggtcag
cgtcctcaccgtcctgcaccaggactggctgaatggcaaggagtacaagt
gcaaggtctccaacaaagccctcccagcccccatcgagaaaaccatctcc
aaagccaaagggcagccccgagaaccacaggtgtacaccctgcccccatc
ccgggaggagatgaccaagaaccaggtcagcctgacctgcctggtcaaag
10 gcttctatcccagcgacatcgccgtggagtgggagagcaatgggcagccg
gagaacaactacgacaccacgcctcccgtgctggactccgacggctcctt
cttccctctatagcgacctcaccgtggacaagagcaggtggcagcagggga
acgtcttctcatgctccgtgatgcaggtctgcacaaccactacacg
cagaagagcctctccctgtctccgggtgcgcgcgacggagaccactgtcc
15 gctcgggcccggcggttgctgccgtctgcacacggtccgcgcgtcgtctgg
aagacctgggctgggccgattgggtgctgtcgccacgggaggtgcaagtg
accatgtgcacgcgtgcccgagccagttccgggcggaacatgca
cgcgcagatcaagacgagcctgcaccgcctgaagcccgcacgggtgccag
cgccctgctgcgtgcccgccagctacaatcccatggtgctcattcaaaag
20 accgacaccggggtgtcgtccagacctatgatgacttgtagccaaaga
ctgccactgcata (SEQ ID NO:180) .

```

In a preferred embodiment, the second polypeptide chain comprises the amino acid sequence of SEQ ID NO:175, which is encoded by the nucleic acid sequence of SEQ ID NO:179.

25 As discussed above, a tetramer is provided comprising two polypeptide chains comprising the sequence of SEQ ID NO:181 and two polypeptide chains comprising the sequence of SEQ ID NO:175.

II.H.20 Dh2CpmFc(-)(N297G)(Y349C)-GDF15(Ndel3):Dh2CpmFc(+)(N297G)(S354C)

30 The designation “Dh2CpmFc(-)(N297G)(Y349C)-GDF15(Ndel3):Dh2CpmFc(+)(N297G)(S354C)” in the instant disclosure refers to a heterodimer comprising (i) a first polypeptide chain comprising a GDF15(Ndel3) polypeptide, the N-terminus of which is linked directly to the C-terminus of a Dh2CpmFc(-)(N297G)(Y349C) polypeptide, and (ii) a second polypeptide chain comprising
35 a Dh2CpmFc(+)(N297G)(S345C) domain. The cysteine clamp mutations allow the first and

second polypeptide chains to be linked via an interchain disulfide bond between C349 of the first polypeptide chain and C354 of the second polypeptide chain.

In certain embodiments, a tetramer is provided, comprising a dimer of two Dh2CpmFc(-)(N297G)(Y349C)-GDF15(Ndel3):Dh2CpmFc(+)(N297G)(S354C)

- 5 heterodimers in which the two first polypeptide chains of each heterodimer are linked via an interchain disulfide bond between their respective GDF15 regions.

More particularly, in a specific embodiment, the tetramer comprises:

(a) two Dh2CpmFc(+)(N297G)(S354C) domains (one each heterodimer) comprising the sequence:

10 PSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVDGVEVHNA
KTKPREEQYGSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPAPIEKTIS
KAKGQPREPQVYITLPPCRKEMTKNQVSLTCLVKGFYPSDIAVEWESNGQP
ENNYKTTTPVLKSDGSFFLYSKLTVDKSRWQQGNVFSVSMHEALHNHYT
QKSLSLSPG (SEQ ID NO:296),

- 15 (b) two Dh2CpmFc(-)(N297G)(Y349C) domains (one each heterodimer) comprising the sequence:

PSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVDGVEVHNA
KTKPREEQYGSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPAPIEKTIS
KAKGQPREPQVCTLPSPREEMTKNQVSLTCLVKGFYPSDIAVEWESNGQP
20 ENNYDTTPVLDSGSDGSFFLYSDLTVDKSRWQQGNVFSVSMHEALHNHYT
QKSLSLSPG (SEQ ID NO:183),

and

(c) two GDF15(Ndel3) polypeptides (one each heterodimer) comprising the sequence of SEQ ID NO:55.

- 25 In a preferred embodiment, the first polypeptide chain comprises the amino acid sequence:

PSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVDGVEVHNA
KTKPREEQYGSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPAPIEKTIS
KAKGQPREPQVCTLPSPREEMTKNQVSLTCLVKGFYPSDIAVEWESNGQP
30 ENNYDTTPVLDSGSDGSFFLYSDLTVDKSRWQQGNVFSVSMHEALHNHYT
QKSLSLSPGGDHCPLGPGRCCRLHTVRASLEDLGWADWVLSPREVQVTMC
IGACPSQFRAANMHAQIKTSLHRLKPDTPAPCCVPASYNPMVLIQKTD
GVSLQTYDDLLAKDCHCI (SEQ ID NO:185),

which is encoded by the nucleic acid sequence:

ccgtcagtccttctcttccccccaaaacccaaggacaccctcatgatctc
 ccggacccctgaggtcacatgcgtggtggtggacgtgagccacgaagacc
 ctgaggtcaagttcaactggtacgtggacggcgtggaggtgcataatgcc
 aagacaaagccgcgggaggagcagtaacggcagcacgtaccgtgtggtcag
 5 cgtcctcacctcctgcaccaggactggctgaatggcaaggagtacaagt
 gcaaggtctccaacaaagccctcccagccccatcgagaaaaccatctcc
 aaagccaaagggcagccccgagaaccacaggtgtgcaccctgccccatc
 ccgggaggagatgaccaagaaccaggtcagcctgacctgcctggtcaaag
 gcttctatcccagcgacatcgccgtggagtgaggagcaatgggcagccg
 10 gagaacaactacgacaccacgcctcccgtgctggactccgacggctcctt
 ctctctctatagcgacctcacctggacaagagcaggtggcagcagggga
 acgtcttctcatgctccgtgatgcatgaggtctgcacaaccactacacg
 cagaagagcctctccctgtctccgggtggagaccactgtccgctcgggcc
 cgggcgttgctgccgtctgcacacgggtccgcgcgtcgctggaagacctgg
 15 gctggggccgattgggtgctgtcgccacgggaggtgcaagtgacctgtgc
 atcgggcgcgtgcccagccagttccggggcggcaaacatgcacgcgcagat
 caagacgagcctgcaccgcctgaagcccgacacgggtgccagcgcctgtc
 gcggtgcccgcagctacaatcccattggtgctcattcaaaagaccgacacc
 ggggtgtcgtccagacctatgatgacttgtagccaaagactgccactg
 20 cata (SEQ ID NO:184).

In a preferred embodiment, the second polypeptide chain comprises the amino acid sequence:

PSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVDGVEVHNA
 KTKPREEQYGSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPAPIEKTIS
 25 KAKGQPREPQVYTLPPCRKEMTKNQVSLTCLVKGFYPSDIAVEWESNGQP
 ENNYKTTTPVLKSDGSFFLYSKLTVDKSRWQQGNVFSVSMHEALHNHYT
 QKSLSLSPGK (SEQ ID NO:182),

which is encoded by the nucleic acid sequence:

ccatccgtgttctctgtttctccaaagccgaaggacaccctgatgatctc
 30 aagaactccggaagtgacttgcgtcgctcggtggacgtgtcacatgaggatc
 cagaggtcaagttcaattggtatgtggacggagtggaagtgcataacgcc
 aagaccaaaccgccgaagaacagtacgggagcacctaccgcgtggtgag
 cgtccttactgtgctccaccaggactggcttaatgggaaggaataacaagt
 gtaaggtgtccaacaaggccctcccgcctcccatcgaaaagaccatctca
 35 aaggcaaaaggggcaaccaagggaacctcaagtgtacaccctgcctccgtg

caggaaggagatgaccaagaaccaggtcagcctgacttgtctcgtgaagg
 gcttctatcccagcgatattgctgtggaatgggagtcaaattggccagccc
 gagaataactacaaaactacccccacccgtgctgaaatctgatgggtcctt
 cttccttttactccaagctgaccgtggacaagagccgctggcaacaaggca
 5 atgtcttttagctgctcagtgatgcatgaggtctccataatcactacact
 cagaagtcactgtccctgtctccgggtaaa (SEQ ID NO:186) .

As discussed above, a tetramer is provided comprising two polypeptide chains comprising the sequence of SEQ ID NO:185 and two polypeptide chains comprising the sequence of SEQ ID NO:182.

10

II.H.21 Dh2CpmFc(-)(N297G)(Y349C)-GDF15(N3D):Dh2CpmFc(+)(N297G)(S354C)

The designation “Dh2CpmFc(-)(N297G)(Y349C)-GDF15(N3D):Dh2CpmFc(+)(N297G)(S354C)” in the instant disclosure refers to a heterodimer comprising (i) a first polypeptide chain comprising a GDF15(N3D) polypeptide,
 15 the N-terminus of which is linked directly to the C-terminus of a Dh2CpmFc(-)(N297G)(Y349C) domain, and (ii) a second polypeptide chain comprising a Dh2CpmFc(+)(N297G)(S345C) domain. The cysteine clamp mutations allow the first and second polypeptide chains to be linked via an interchain disulfide bond between C349 of the first polypeptide and C354 of the second polypeptide.

20 In certain embodiments, a tetramer is provided, comprising a dimer of two Dh2CpmFc(-)(N297G)(Y349C)-GDF15(N3D):Dh2CpmFc(+)(N297G)(S354C) heterodimers in which the two first polypeptide chains of each heterodimer are linked via an interchain disulfide bond between their respective GDF15 regions.

More particularly, in a specific embodiment, the tetramer comprises:

25 (a) two Dh2CpmFc(+)(N297G)(S354C) domains (one each heterodimer) comprising the sequence of SEQ ID NO:296,

(b) two Dh2CpmFc(-)(N297G)(Y349C) domains (one each heterodimer) comprising the sequence of SEQ ID NO:183, and

(c) two GDF15(N3D) polypeptide (one each heterodimer) comprising the sequence of
 30 SEQ ID NO:52.

In a preferred embodiment, the first polypeptide chain comprises the amino acid sequence:

PSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVDGVEVHNA
 KTKPREEQYGSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPAPIEKTIS
 35 KAKGQPREPQVCTLPSPREEMTKNQVSLTCLVKGFYPSDIAVEWESNGQP
 ENNYDTTPPVLDSDGSFFLYSDLTVDKSRWQQGNVFSQSVMEALHNHYT

QKSLSLSPGARDGDHCPGPRCCRLHTVRASLEDLGWADWVLSPREVQV
 TMCIGACPSQFRAANMHAQIKTSLHRLKPDTPAPCCVPASYNPMVLIQK
 TDTGVSLQTYDDLAKDCHCI (SEQ ID NO:188),

which is encoded by the nucleic acid sequence:

5 ccgtcagtccttctcttccccccaaaacccaaggacaccctcatgatctc
 ccggacccctgaggtcacatgcgtgggtggacgtgagccacgaagacc
 ctgaggtcaagttcaactggtagcgtggacggcgtggaggtgcataatgcc
 aagacaaagccgcgggaggagcagtagcgggagcacgtaccgtgtggtcag
 cgtcctcaccgtcctgcaccaggactggctgaatggcaaggagtacaagt
 10 gcaaggtctccaacaaagccctcccagcccccatcgagaaaaccatctcc
 aaagccaaagggcagccccgagaaccacaggtgtgaccctgcccccatc
 ccgggaggagatgaccaagaaccaggtcagcctgacctgcctggtcaaag
 gcttctatcccagcgacatcgccgtggagtgggagagcaatgggcagccg
 gagaacaactacgacaccacgcctcccgtgctggactccgacggctcctt
 15 ctctctctatagcgacctcaccgtggacaagagcaggtggcagcagggga
 acgtcttctcatgctccgtgatgcatgaggtctctgcacaaccactacacg
 cagaagagcctctccctgtctccgggtgcgcgcgacggagaccactgtcc
 gctcgggccccggcggttgctgcccgtctgcacacgggtccgcgcgtcgctgg
 aagacctgggctgggcccattgggtgctgtcgccacgggaggtgcaagtg
 20 accatgtgcatcggcgcgtgcccagagccagttccgggcccgaacatgca
 cgcgagatcaagacgagcctgcaccgcctgaagcccgcacacgggtgccag
 cgccctgctgctgcccgcagctacaatcccatgggtgctcattcaaaag
 accgacaccggggtgtcgtccagacctatgatgacttgtagccaaaga
 ctgccactgcata (SEQ ID NO:187).

25 In a preferred embodiment, the second polypeptide chain comprises the amino acid sequence of SEQ ID NO:182, which is encoded by the nucleic acid sequence of SEQ ID NO:186.

As discussed above, a tetramer is provided comprising two polypeptide chains comprising the sequence of SEQ ID NO:188 and two polypeptide chains comprising the
 30 sequence of SEQ ID NO:182.

II.H.22 Dh2CpmFc(-)(N297G)(A287C)-GDF15(Ndel3):Dh2CpmFc(+)(N297G)(L306C)

The designation “Dh2CpmFc(-)(N297G)(A287C)-GDF15(Ndel3):Dh2CpmFc(+)(N297G)(L306C)” in the instant disclosure refers to a
 35 heterodimer comprising (i) a first polypeptide chain comprising GDF15(Ndel3) polypeptide,

the N-terminus of which is linked directly to the C-terminus of a Dh2CpmFc(-)(N297G)(A287C) domain, and (ii) a second polypeptide chain comprising a Dh2CpmFc(+)(N297G)(L306C) domain. The cysteine clamp mutations allow the first and second polypeptide chains to be linked via an interchain disulfide bond between C287 of the first monomer and C306 of the second monomer.

In certain embodiments, a tetramer is provided, comprising a dimer of two Dh2CpmFc(-)(N297G)(A287C)-GDF15(Nde13):Dh2CpmFc(+)(N297G)(L306C) heterodimers in which the two first polypeptide chains of each heterodimer are linked via an interchain disulfide bond between their respective GDF15 regions.

More particularly, in a specific embodiment, a tetramer is provided which comprises:
(a) two Dh2CpmFc(+)(N297G)(L306C) domains (one each heterodimer) comprising the sequence:

PSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVDGVEVHNC
KTKPREEQYGSTYRVVSVCTV LHQDWLNGKEYKCKVSNKALPAPIEKTIS
KAKGQPREPQVYTLPPSRKEMTKNQVSLTCLVKGFYPSDIAVEWESNGQP
ENNYKTTTPVLKSDGSFFLYSKLTVDKSRWQQGNVFSCSVMHEALHNHYT
QKSLSLSPG (SEQ ID NO:297),

(b) two Dh2CpmFc(-)(N297G)(A287C) (one each heterodimer) comprising the sequence:

PSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVDGVEVHNC
KTKPREEQYGSTYRVVSVCTV LHQDWLNGKEYKCKVSNKALPAPIEKTIS
KAKGQPREPQVYTLPPSREEMTKNQVSLTCLVKGFYPSDIAVEWESNGQP
ENNYDTTPVLDSDGSFFLYSDLTVDKSRWQQGNVFSCSVMHEALHNHYT
QKSLSLSPG (SEQ ID NO:192),

and

(c) two GDF15(Nde13) polypeptides (one each heterodimer) comprising the sequence of SEQ ID NO:55.

In a preferred embodiment, the first polypeptide chain comprises the amino acid sequence:

PSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVDGVEVHNC
KTKPREEQYGSTYRVVSVCTV LHQDWLNGKEYKCKVSNKALPAPIEKTIS
KAKGQPREPQVYTLPPSREEMTKNQVSLTCLVKGFYPSDIAVEWESNGQP
ENNYDTTPVLDSDGSFFLYSDLTVDKSRWQQGNVFSCSVMHEALHNHYT
QKSLSLSPGGDHCPLGPGRCCRLHTVRASLEDLGWADWVLSPREVQVTMC

IGACPSQFRAANMHAQIKTSLHRLKPDTPAPCCVPASYNPMVLIQKTD
 GVSLQTYDDLLAKDCHCI (SEQ ID NO:194),

which is encoded by the nucleic acid sequence:

ccgtcagtccttcttccccccaaaacccaaggacaccctca
 5 tgatctcccgagccctgaggtcacatgctggtggtggacgtgagccac
 gaagaccctgaggtcaagttcaactggtacgtggacggcgtggaggtgca
 taattgcaagacaaagccgaggagcagtagcggcagcacgtaccgtg
 tggtcagcgtctgcaccgtcctgcaccaggactggtgaatggcaaggag
 tacaagtgaaggtctccaacaaagccctccagcccccatcgaaaaac
 10 catctccaaagccaaagggcagccccgagaaccacaggtgtacaccctgc
 ccccatcccgaggagatgaccaagaaccaggtcagcctgacctgcctg
 gtcaaaggcttctatcccagcgacatcgccgtggagtgggagagcaatgg
 gcagccggagaacaactacgacaccacgcctcccggtgctggactccgacg
 gctccttcttctctatagcgacctcaccgtggacaagagcaggtggcag
 15 caggggaacgtcttctcatgctccgtgatgcatgaggctctgcacaacca
 ctacacgcagaagagcctctccctgtctccgggtggagaccactgtccgc
 tcgggccccggcgcttgctgccgtctgcacacggtcgcgcgctcgctggaa
 gacctgggctgggcccattgggtgctgtgccacgggaggtgcaagtga
 catgtgcatcggcgctgcccagccagttccgggcccgaacatgcaag
 20 cgcagatcaagacgagcctgcaccgcctgaagcccagacaggtgccagcg
 cctgctgctgcccgcagctacaatcccatggtgctcattcaaaagac
 cgacaccggggtgctgctccagacctatgatgacttgtagccaaagact
 gccactgcata (SEQ ID NO:193) .

25 In a preferred embodiment, the second polypeptide chain comprises the amino acid
 sequence:

PSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVDGVEVHNC
 KTKPREEQYGSTYRVVSVCTV LHQDWLNGKEYKCKVSNKALPAPIEKTIS
 KAKGQPREPQVYTLPPSRKEMTKNQVSLTCLVKGFYPSDIAVEWESNGQP
 ENNYKTTTPVLKSDGSFFLYSKLTVDKSRWQQGNVFSVSMHEALHNHYT
 30 QKSLSLSPGK (SEQ ID NO:191) ,

which is encoded by the nucleic acid sequence:

ccgtcagtccttcttccccccaaaacccaaggacaccctca
 tgatctcccgagccctgaggtcacatgctggtggtggacgtgagccac
 gaagaccctgaggtcaagttcaactggtacgtggacggcgtggaggtgca

taattgcaagacaaagccgcgggaggagcagtacggcagcacgtaccgtg
 tggtcagcgtctgcaccgtcctgcaccaggactggctgaatggcaaggag
 tacaagtgcaaggtctccaacaaagccctcccagcccccatcgagaaaac
 catctccaaagccaaagggcagccccgagaaccacaggtgtacacctgc
 5 ccccatcccggaaggagatgaccaagaaccaggtcagcctgacctgacctg
 gtcaaaggcttctatcccagcgacatcgccgtggagtgggagagcaatgg
 gcagccggagaacaactacaagaccacgcctcccgtgctgaagtccgacg
 gctccttcttctctatagcaagctcaccgtggacaagagcaggtggcag
 caggggaacgtcttctcatgctccgtgatgcatgaggctctgcacaacca
 10 ctacacgcagaagagcctctccctgtctccgggtaaa (SEQ ID NO:195) .

As discussed above, a tetramer is provided comprising two polypeptide chains comprising the sequence of SEQ ID NO:194 and two polypeptide chains comprising the sequence of SEQ ID NO:191.

15 *II.H.23 Dh2CpmFc(-)(N297G)(A287C)-GDF15(N3D):Dh2CpmFc(+)(N297G)(L306C)*

The designation “Dh2CpmFc(-)(N297G)(A287C)-
 GDF15(N3D):Dh2CpmFc(+)(N297G)(L306C)” in the instant disclosure refers to a
 heterodimer comprising (i) a first polypeptide chain comprising a GDF15(N3D) polypeptide,
 the N-terminus of which is linked directly to the C-terminus of a
 20 Dh2CpmFc(-)(N297G)(A287C) domain, and (ii) a second polypeptide chain comprising a
 Dh2CpmFc(+)(N297G)(L306C) domain. The cysteine clamp mutations allow the first and
 second polypeptide chains to be linked via an interchain disulfide bond between C287 of the
 first polypeptide chain and C306 of the second polypeptide chain.

In certain embodiments, a tetramer is provided, comprising a dimer of two
 25 Dh2CpmFc(-)(N297G)(A287C)-GDF15(N3D):Dh2CpmFc(+)(N297G)(L306C) heterodimers
 in which the two first polypeptide chains of each heterodimer are linked via an interchain
 disulfide bond between their respective GDF15 regions.

More particularly, in a specific embodiment, a tetramer is provided which comprises:

- 30 (a) two Dh2CpmFc(+)(N297G)(L306C) domains (one each heterodimer) comprising
 the sequence of SEQ ID NO:297,
- (b) two Dh2CpmFc(-)(N297G)(A287C) domains (one each heterodimer) comprising
 the sequence of SEQ ID NO:192, and
- (c) two GDF15(N3D) polypeptides (one each heterodimer) comprising the sequence
 of SEQ ID NO:52.

35 In a preferred embodiment, the first polypeptide chain comprises the amino acid
 sequence:

PSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVDGVEVHNC
 KTKPREEQYGSTYRVVSVCTV LHQDWLNGKEYKCKVSNKALPAPIEKTIS
 KAKGQPREPQVYTLPPSREEMTKNQVSLTCLVKGFYPSDIAVEWESNGQP
 ENNYDTTPPVLDSDGSFFLYSDLTVDKSRWQQGNVSCSVMHEALHNHYT
 5 QKSLSLSPGARDGDHCPLGPGRCRLHTVRASLEDLGWADWVLSPREVQV
 TMCIGACPSQFRAANMHAQIKTSLHRLKPDTPAPCCVPASYNPMVLIQK
 TDTGVSLQTYDDLLAKDCHCI (SEQ ID NO:197),

which is encoded by the nucleic acid sequence:

ccgtcagtccttctcttccccccaaaacccaaggacaccctca
 10 tgatctcccgaccctgaggtcacatgcgtggtggacgtgagccac
 gaagaccctgaggtcaagttcaactggtacgtggacggcgtggaggtgca
 taattgcaagacaaagccgcgggaggagcagtagcgcagcacgtaccgtg
 tggtcagcgtctgcaccgtcctgcaccaggactggctgaatggcaaggag
 tacaagtgcaaggtctccaacaaagccctcccagcccccatcgagaaaac
 15 catctccaaagccaaagggcagccccgagaaccacaggtgtacaccctgc
 ccccatcccgaggagatgaccaagaaccaggtcagcctgacctgcctg
 gtcaaaggcttctatcccagcgacatcgccgtggagtgggagagcaatgg
 gcagccggagaacaactacgacaccacgcctcccgctgctggactccgacg
 gctccttcttctctatagcgacctcaccgtggacaagagcaggtggcag
 20 caggggaacgtcttctcatgctcctgatgcatgaggctctgcacaacca
 ctacacgcagaagagcctctccctgtctccgggtgcgcgcgacggagacc
 actgtccgctcgggccccggcggttctgctgccgtctgcacacggtccgcgcg
 tcgctggaagacctgggctgggcccattgggtgctgtcgccacgggaggt
 gcaagtgacctatgtgcatcggcgcgtgcccagagccagttccggggcgga
 25 acatgcacgcgcagatcaagacgagcctgcaccgcctgaagcccgacacg
 gtgccagcgccctgctgcgtgcccgcagctacaatcccatggtgctcat
 tcaaaagaccgacaccggggtgtcgctccagacctatgatgacttgtag
 ccaaagactgccactgcata (SEQ ID NO:196).

In a preferred embodiment, the second polypeptide chain comprises the amino acid
 30 sequence of SEQ ID NO:191, which is encoded by the nucleic acid sequence of SEQ ID
 NO:195.

As discussed above, a tetramer is provided comprising two polypeptide chains
 comprising the sequence of SEQ ID NO:197 and two polypeptide chains comprising the
 sequence of SEQ ID NO:191.

35

*II.H.24 Dh2CpmFc(-)(N297G)(A287C)(Y349C)-GDF15(Ndel3):
Dh2CpmFc(+)(N297G)(L306C)(S354C)*

The designation “Dh2CpmFc(-)(N297G)(A287C)(Y349C)-GDF15(Ndel3):Dh2CpmFc(+)(N297G)(L306C)(S354C)” in the instant disclosure refers to a heterodimer comprising (i) a first polypeptide chain comprising GDF15(Ndel3) polypeptide, the N-terminus of which is linked directly to the C-terminus of a Dh2CpmFc(-)(N297G)(A287C)(Y349C) domain, and (ii) a second polypeptide chain comprising a Dh2CpmFc(+)(N297G)(L306C)(S354C) domain. The cysteine clamp mutations allow the first and second polypeptide chains to be linked via an interchain disulfide bond between C287 of the first polypeptide chain and C306 of the second polypeptide chain and between the C349 of the first polypeptide chain and C349 of the second polypeptide chain.

In certain embodiments, a tetramer is provided, comprising a dimer of two Dh2CpmFc(-)(N297G)(A287C)(Y349C)-GDF15(Ndel3):Dh2CpmFc(+)(N297G)(L306C)(S354C) heterodimers in which the two first polypeptide chains of each heterodimer are linked via an interchain disulfide bond between their respective GDF15 regions.

More particularly, in a specific embodiment, a tetramer is provided which comprises:

(a) two Dh2CpmFc(+)(N297G)(L306C)(S354C) domains (one each heterodimer) comprising the sequence:

PSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVDGVEVHNC
KTKPREEQYGSTYRVVSVCTV LHQDWLNGKEYKCKVSNKALPAPIEKTIS
KAKGQPREPQVYTLPPCRKEMTKNQVSLTCLVKGFYPSDIAVEWESNGQP
ENNYKTTTPVLKSDGSFFLYSKLTVDKSRWQQGNVFSCSVMHEALHNHYT
QKSLSLSPG (SEQ ID NO:298),

(b) two Dh2CpmFc(-)(N297G)(A287C)(Y349C) domains (one each heterodimer) comprising the sequence:

PSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVDGVEVHNC
KTKPREEQYGSTYRVVSVCTV LHQDWLNGKEYKCKVSNKALPAPIEKTIS
KAKGQPREPQVCTLPPSREEMTKNQVSLTCLVKGFYPSDIAVEWESNGQP
ENNYDTPPVLDSDGSFFLYSDLTVDKSRWQQGNVFSCSVMHEALHNHYT
QKSLSLSPG (SEQ ID NO:199),

and

(c) two GDF15(Ndel3) polypeptides (one each heterodimer) comprising the sequence of SEQ ID NO:55.

In a preferred embodiment, the first polypeptide chain comprises the amino acid sequence:

PSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVDGVEVHNC
 KTKPREEQYGSTYRVVSVCTV LHQDWLNGKEYKCKVSNKALPAPIEKTIS
 5 KAKGQPREPQVCTLPSPREEMTKNQVSLTCLVKGFYPSDIAVEWESNGQP
 ENNYDTTPPVLDSDGSFFLYSDLTVDKSRWQQGNVFSQSVMEALHNHYT
 QKSLSLSPGGDHCPLGPGRCCRLHTVRASLEDLGWADWVLSPREVQVTMC
 IGACPSQFRAANMHAQIKTSLHRLKPDTPVAPCCVPASYNPMVLIQKTD
 GVSLQTYDDLAKDCHCI (SEQ ID NO:201),

10 which is encoded by the nucleic acid sequence:

ccgtcagtccttctcttccccccaaaacccaaggacaccctca
 tgatctcccgaccctgaggtcacatgcgtggtggtgacgtgagccac
 gaagaccctgaggtcaagttcaactggtacgtggacggcgtggaggtgca
 taattgcaagacaaagccgcgaggagcagtagcggcagcacgtaccgtg
 15 tggtcagcgtctgcaccgtcctgcaccaggactggctgaatggcaaggag
 tacaagtgcaggtctccaacaaagccctccagcccccatcgagaaaac
 catctccaaagccaaagggcagccccgagaaccacaggtgtgcaccctgc
 ccccatcccgaggagatgaccaagaaccaggtcagcctgacctgctg
 gtcaaaggcttctatcccagcgacatcgccgtggagtgggagagcaatgg
 20 gcagccggagaacaactacgacaccacgcctcccggtgctggactccgacg
 gctccttcttctctatagcgacctcaccgtggacaagagcaggtggcag
 caggggaacgtcttctcatgctccgtgatgcatgaggtctgcacaacca
 ctacacgcagaagagcctctccctgtctccgggtggagaccactgtccgc
 tcgggccccggcggttgcgtgcccgtctgcacacggtccgcgcgtcgctggaa
 25 gacctgggctgggcccattgggtgctgtgccacgggaggtgcaagtgc
 catgtgcacatcggcgcgtgcccagccagttccgggcggaacatgcacg
 cgcagatcaagacgagcctgcaccgcctgaagcccgcacacggtgccagcg
 ccctgctgcgtgcccgccagctacaatcccatggtgctcattcaaaagac
 cgacaccggggtgctgctccagacctatgatgacttgtagccaaagact
 30 gccactgcata (SEQ ID NO:200).

In a preferred embodiment, the second polypeptide chain comprises the amino acid sequence:

PSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVDGVEVHNC
 KTKPREEQYGSTYRVVSVCTV LHQDWLNGKEYKCKVSNKALPAPIEKTIS

KAKGQPREPQVYTLPPCRKEMTKNQVSLTCLVKGFYPSDIAVEWESNGQP
ENNYKTTTPVLKSDGSFFLYSKLTVDKSRWQQGNVFSVSMHEALHNHYT
QKSLSLSPGK (SEQ ID NO:198),

which is encoded by the nucleic acid sequence:

5 ccgtcagtccttctcttccccccaaaacccaaggacaccctca
tgatctcccgaccctgaggtcacatgcgtggtggtggacgtgagccac
gaagaccctgaggtcaagttcaactggtacgtggacggcgtggaggtgca
taattgcaagacaaagccgcgggaggagcagtagcggcagcacgtaccgtg
tggtcagcgtctgcaccgtcctgcaccaggactggctgaatggcaaggag
10 tacaagtgaaggtctccaacaaagccctcccagcccccatcgagaaaac
catctccaaagccaaagggcagccccgagaaccacaggtgtacaccctgc
ccccatgccggaaggagatgaccaagaaccaggtcagcctgacctgcctg
gtcaaaggcttctatcccagcgacatcgccgtggagtgggagagcaatgg
gcagccggagaacaactacaagaccacgcctcccgtgctgaagtccgacg
15 gctccttcttctctatagcaagctcaccgtggacaagagcaggtggcag
caggggaacgtcttctcatgctccgtgatgcatgaggctctgcacaacca
ctacacgcagaagagcctctccctgtctccgggtaaa (SEQ ID NO:202).

As discussed above, a tetramer is provided comprising two polypeptide chains
comprising the sequence of SEQ ID NO:201 and two polypeptide chains comprising the
20 sequence of SEQ ID NO:198.

*II.H.25 Dh2CpmFc(-)(N297G)(A287C)(Y349C)-GDF15(N3D):
Dh2CpmFc(+)(N297G)(L306C)(S354C)*

The designation
25 “Dh2CpmFc(-)(N297G)(A287C)(Y349C)-GDF15(N3D):Dh2CpmFc(+)(N297G)(L306C)(S354C)” in the instant disclosure refers to a heterodimer comprising (i) a first monomer comprising GDF15(N3D) polypeptide, the N-terminus of which is linked directly to the C-terminus of a Dh2CpmFc(-)(N297G)(A287C)(Y349C) domain, and (ii) a second polypeptide chain comprising a Dh2CpmFc(+)(N297G)(L306C)(S354C) domain. The cysteine clamp
30 mutations allow the first and second polypeptide chains to be linked via an interchain disulfide bond between C287 of the first polypeptide chain and C306 of the second polypeptide chain and between the C349 of the first polypeptide chain and C349 of the second polypeptide chain.

In certain embodiments, a tetramer is provided, comprising a dimer of two
35 Dh2CpmFc(-)(N297G)(A287C)(Y349C)-GDF15(N3D):Dh2CpmFc(+)(N297G)(L306C)(S354C)

4C) heterodimers in which the two first polypeptide chains of each heterodimer are linked via an interchain disulfide bond between their respective GDF15 regions.

More particularly, in a specific embodiment, a tetramer is provided which comprises:

- (a) two Dh2CpmFc(+)(N297G)(L306C)(S354C) domains (one each heterodimer) comprising the sequence of SEQ ID NO:298,
- (b) two Dh2CpmFc(-)(N297G)(A287C)(Y349C) domains (one each heterodimer) comprising the sequence of SEQ ID NO:199, and
- (c) two GDF15(N3D) polypeptides (one each heterodimer) comprising the sequence of (SEQ ID NO:52).

- 10 In a preferred embodiment, the first polypeptide chain comprises the amino acid sequence:

PSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVDGVEVHNC
 KTKPREEQYGSTYRVVSVCTV LHQDWLNGKEYKCKVSNKALPAPIEKTIS
 KAKGQPREPQVCTLPSPREEMTKNQVSLTCLVKGFYPSDIAVEWESNGQP
 15 ENNYDTTPPVLDSDGSFFLYSDLTVDKSRWQQGNV FSCSVMHEALHNHYT
 QKSLSLSPGARDGDHCP LGPGRCCRLHTVRASLEDLGWADWVLSPREVQV
 TMCIGACPSQFRAANMHAQIKTSLHRLKPDTPVAPCCVPASYNPMVLIQK
 TDTGVSLQTYDDLAKDCHCI (SEQ ID NO:204),

which is encoded by the nucleic acid sequence:

- 20 ccgtcagtccttctcttccccccaaaacccaaggacaccctca
 tgatctcccgaccctgaggtcacatgcgtggtggacgtgagccac
 gaagaccctgaggtcaagttcaactggtacgtggacggcgtggaggtgca
 taattgcaagacaaagccgcgggaggagcagtacggcagcacgtaccgtg
 tggtcagcgtctgcaccgtcctgcaccaggactggctgaatggcaaggag
 25 tacaagtgcaaggtctccaacaaagccctcccagcccccatcgagaaaac
 catctccaaagccaaagggcagccccgagaaccacaggtgtgcaccctgc
 ccccatcccgggaggagatgaccaagaaccaggtcagcctgacctgctg
 gtcaaaggcttctatcccagcgacatcgccgtggagtgggagagcaatgg
 gcagccggagaacaactacgacaccacgcctcccgtgctggactccgacg
 30 gctccttcttctctatagcgacctcaccgtggacaagagcaggtggcag
 caggggaacgtcttctcatgctccgtgatgcatgaggctctgcacaacca
 ctacacgcagaagagcctctccctgtctccgggtgcgcgacggagacc
 actgtccgctcgggccccggcggttgctgccgtctgcacacggtccgcg
 tcgctggaagacctgggctgggcccattgggtgctgtcgccacgggaggt
 35 gcaagtgaccatgtgcatcggcgcgtgcccagaccagttccgggcgga

acatgcacgcgcagatcaagacgagcctgcaccgcctgaagcccgacacg
gtgccagcgcacctgctgcgtgcccgcagctacaatcccatggtgctcat
tcaaaagaccgacaccggggtgctcgtccagacctatgatgacttggttag
ccaaagactgccactgcata (SEQ ID NO:203) .

5 In a preferred embodiment, the second polypeptide chain comprises the amino acid sequence of SEQ ID NO:198, which is encoded by the nucleic acid sequence of SEQ ID NO:202.

As discussed above, a tetramer is provided comprising two polypeptide chains comprising the sequence of SEQ ID NO:204 and two polypeptide chains comprising the
10 sequence of SEQ ID NO:198.

II.H.26 GG-Dh2CpmFc(-)-GDF15(Ndel3):GG-Dh2CpmFc(+)

The designation “GG- Dh2CpmFc(-)-GDF15(Ndel3):GG-Dh2CpmFc(+)” in the instant disclosure refers to a heterodimer comprising (i) a first polypeptide chain comprising a
15 GDF15(Ndel3) polypeptide, the N-terminus of which is linked directly to the C-terminus of a GG-Dh2CpmFc(-) domain, and (ii) a second polypeptide chain comprising a GG-Dh2CpmFc(+) domain.

In certain embodiments, a tetramer is provided, comprising a dimer of two GG-Dh2CpmFc(-)-GDF15(Ndel3):GG-Dh2CpmFc(+) heterodimers in which the two first
20 polypeptide chains of each heterodimer are linked via an interchain disulfide bond between their respective GDF15 regions.

More particularly, in a specific embodiment, the tetramer comprises:

(a) two GG-Dh2CpmFc(+) domains (one each heterodimer) comprising the sequence:

GGPSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVDGVEVH
25 NAKTKPREEQYNSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPAPIEKT
ISKAKGQPREPQVYTLPPSRKEMTKNQVSLTCLVKGFYPSDIAVEWESNG
QPENNYKTTTPVLKSDGSFFLYSKLTVDKSRWQQGNVFSCSVMHEALHNH
YTQKSLSLSPG (SEQ ID NO:299) ,

(b) two GG-Dh2CpmFc(-) domains (one each heterodimer) comprising the sequence:

30 GGPSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVDGVEVH
NAKTKPREEQYNSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPAPIEKT
ISKAKGQPREPQVYTLPPSREEMTKNQVSLTCLVKGFYPSDIAVEWESNG
QPENNYDTTPVLDSGFFLYSDLTVDKSRWQQGNVFSCSVMHEALHNH
YTQKSLSLSPG (SEQ ID NO:206) ,

35 and

(c) two GDF15(NdeI3) polypeptides (one each heterodimer) comprising the sequence of SEQ ID NO:55.

In a preferred embodiment, the first polypeptide chain comprises the amino acid sequence:

5 GGPSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVDGVEVH
NAKTKPREEQYNSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPAPIEKT
ISKAKGQPREPQVYTLPPSREEMTKNQVSLTCLVKGFYPSDIAVEWESNG
QPENNYDTTPPVLDSDGSFFLYSDLTVDKSRWQQGNVFSQSVMEALHNH
YTQKSLSLSPGGDHCPLGPGRCRLHTVRASLEDLGWADWVLSPREVQVT
10 MCIGACPSQFRAANMHAQIKTSLHRLKPDTPAPCCVPASYNPMVLIQKT
DTGVSLQTYDDLAKDCHCI (SEQ ID NO:208),

which is encoded by the nucleic acid sequence:

ggtagggccgtagctcttctcttccccccaaaaccaaggaca
ccctcatgatctcccgagccctgaggtcacatgctgggtggacgtg
15 agccacgaagacctgaggtcaagttcaactggtacgtggacggcgtgga
ggtagcataatgccaaagacaaagccgagggaggagcagtacaacagcagct
acctgtggtcagcgtcctcaccgtcctgcaccaggactggctgaatggc
aaggagtacaagtgaaggtctccaacaaagccctccagcccccatcga
gaaaaccatctccaaagccaaagggcagccccgagaaccacaggtgtaca
20 ccctgcccccatcccgaggagatgaccaagaaccaggtcagcctgacc
tgcttggtcaaaggcttctatcccgagcagatcgccgtggagtgggagag
caatgggcagccggagaacaactacgacaccacgcctcccgtagctggact
ccgacggctccttcttctctatagcgacctaccgtggacaagagcagg
tggcagcaggggaacgtcttctcatgctccgtgatgcatgaggctctgca
25 caaccactacacgcagaagagcctctccctgtctccgggtggagaccact
gtccgctcgggcccggcggttgctgcctgtctgcacacgggtccgcgcgtcg
ctggaagacctgggctgggcccattgggtgctgctgccacgggaggtgca
agtgacctatgtgcatcggcgctgcccagaccagttccgggaggcaaca
tgcaacgcgcagatcaagacgagcctgcaccgcctgaagcccagacaggtg
30 ccagcgcctgtgctgctgcccgcagctacaatcccatggtgctcattca
aaagaccgacaccggggtgctgctccagacctatgatgacttggttagcca
aagactgccactgcata (SEQ ID NO:207).

In an embodiment employing the VH21 signal sequence, in a preferred embodiment, the first polypeptide chain comprises the amino acid sequence (signal sequence single
35 underlined):

MEWSWVFLFFLSVTTGVHSGGPSVFLFPPKPKDTLMISRTPEVTCVVVDV
 SHEDPEVKFNWYVDGVEVHNAKTKPREEQYNSTYRVVSVLTVLHQDWLNG
 KEYKCKVSNKALPAPIEKTISKAKGQPREPQVYTLPPSREEMTKNQVSLT
 CLVKGFYPSDIAVEWESNGQPENNYDTPPVLDSDGSFFLYSDLTVDKSR
 5 WQQGNVFSCSVMHEALHNHYTQKSLSLSPGGDHCPLGPGRCCRLHTVRAS
 LEDLGWADWVLSPREVQVTMCIGACPSQFRAANMHAQIKTSLHRLKPDVT
 PAPCCVPASYNPMVLIQKTDGTGVSLOTYDDLAKDCHCI (SEQ ID NO:243),

which is encoded by the nucleic acid sequence (signal sequence underlined):

atggaatggagctgggtctttctcttcttctcctgtcagtaacgactgggtgt
 10 ccactccggtggcccgtcagtccttctcttcccccaaaaccaaggaca
 ccctcatgatctcccgaccctgaggtcacatgctggtggtggacgtg
 agccacgaagaccctgaggtcaagttcaactggtacgtggacggcgtgga
 ggtgcataatgccaaagacaaagccgcgggaggagcagtacaacagcacgt
 accgtgtggtcagcgtcctcaccgtcctgcaccaggactggctgaatggc
 15 aaggagtacaagtgaaggtctccaacaaagccctcccagcccccatcga
 gaaaaccatctccaaagccaaagggcagccccgagaaccacaggtgtaca
 ccctgcccccatcccgaggagatgaccaagaaccaggtcagcctgacc
 tgcttgggtcaaaggcttctatcccagcgacatcgccgtggagtgggagag
 caatgggcagccggagaacaactacgacaccacgcctcccggtgctggact
 20 ccgacggctccttcttctctatagcgacctcaccgtggacaagagcagg
 tggcagcaggggaacgtcttctcatgctccgtgatgcatgaggctctgca
 caaccactacacgcagaagagcctctccctgtctccgggtggagaccact
 gtccgctcgggccccggcggttgctgccgtctgcacacggtccgcgcgtcg
 ctggaagacctgggctgggcccattgggtgctgtcgccacgggaggtgca
 25 agtgaccatgtgcatcggcgcgtgcccagaccagttccgggcccgaaca
 tgcacgcgcagatcaagacgagcctgcaccgcctgaagcccagacagggtg
 ccagcgccctgctgctgctgcccgcagctacaatcccatggtgctcattca
 aaagaccgacaccggggtgctgctccagacctatgatgacttgtagcca
 aagactgccactgcata (SEQ ID NO:244).

30 In a preferred embodiment, the second polypeptide chain comprises the amino acid sequence:

GGPSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVDGVEVH
 NAKTKPREEQYNSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPAPIEKT
 ISKAKGQPREPQVYTLPPSRKEMTKNQVSLTCLVKGFYPSDIAVEWESNG

QPENNYKTPPVLKSDGSFFLYSKLTVDKSRWQQGNVFSCSVMHEALHNNH
YTQKSLSLSPGK (SEQ ID NO:205),

which is encoded by the nucleic acid sequence:

ggtggcccgctcagtccttctcttccccccaaaacccaaggaca
5 ccctcatgatctcccgagccctgaggtcacatgctggtggtggacgtg
agccacgaagaccctgaggtcaagttcaactggtacgtggacggcgtgga
ggtgcataatgccaaagacaaagccgcgggaggagcagtacaacagcacgt
accgtgtggtcagcgtcctcaccgtcctgcaccaggactggctgaatggc
aaggagtacaagtgaaggtctccaacaaagccctcccagcccccatcga
10 gaaaaccatctccaaagccaaagggcagccccgagaaccacaggtgtaca
ccctgcccccatcccgaaggagatgaccaagaaccaggtcagcctgacc
tgcttgggtcaaaggcttctatcccagcgacatcgccgtggagtgggagag
caatgggcagccggagaacaactacaagaccacgcctcccgtgctgaagt
ccgacggctccttcttctctatagcaagctcaccgtggacaagagcagg
15 tggcagcaggggaacgtcttctcatgctccgtgatgcatgaggctctgca
caaccactacacgcagaagagcctctccctgtctccgggtaaa (SEQ ID NO:209).

In an embodiment employing the VH21 signal sequence, in a preferred embodiment, the second polypeptide chain comprises the amino acid sequence (signal sequence single underlined):

20 MEWSWVFLFFLSVTTGVHSGGPSVFLFPPKPKDTLMISRTPEVTCVVVDV
SHEDPEVKFNWYVDGVEVHNAKTKPREEQYNSTYRVVSVLTVLHQDWLNG
KEYKCKVSNKALPAPIEKTISKAKGQPREPQVYTLPPSRKEMTKNQVSLT
CLVKGFYPSDIAVEWESNGQPENNYKTPPVLKSDGSFFLYSKLTVDKSR
WQQGNVFSCSVMHEALHNNHYTQKSLSLSPGK (SEQ ID NO:245),

25 which is encoded by the nucleic acid sequence (signal sequence underlined):

atggaatggagctgggtctttctcttcttctcctgtcagtaacgactgggtgt
ccactccgggtggcccgctcagtccttcttccccccaaaacccaaggaca
ccctcatgatctcccgagccctgaggtcacatgctggtggtggacgtg
agccacgaagaccctgaggtcaagttcaactggtacgtggacggcgtgga
30 ggtgcataatgccaaagacaaagccgcgggaggagcagtacaacagcacgt
accgtgtggtcagcgtcctcaccgtcctgcaccaggactggctgaatggc
aaggagtacaagtgaaggtctccaacaaagccctcccagcccccatcga
gaaaaccatctccaaagccaaagggcagccccgagaaccacaggtgtaca
ccctgcccccatcccgaaggagatgaccaagaaccaggtcagcctgacc

tgccctgggtcaaaggcttctatcccagcgacatcgccgtggagtgaggagag
 caatgggcagccggagaacaactacaagaccacgcctcccgtgctgaagt
 ccgacggctccttcttctctatagcaagctcaccgtggacaagagcagg
 tggcagcaggggaacgtcttctcatgctccgtgatgcatgaggctctgca
 5 caaccactacacgcagaagagcctctccctgtctccgggtaaa (SEQ ID NO:246) .

As discussed above, a tetramer is provided comprising two polypeptide chains comprising the sequence of SEQ ID NO:208 and two polypeptide chains comprising the sequence of SEQ ID NO:205.

10 *II.H.27 GG-Dh2CpmFc(-)-GDF15(N3D):GG-Dh2CpmFc(+)*

The designation “GG-Dh2CpmFc(-)-GDF15(N3D):GG-Dh2CpmFc(+)” in the instant disclosure refers to a heterodimer comprising (i) a first polypeptide chain comprising a GDF15(N3D) polypeptide, the N-terminus of which is linked directly to the C-terminus of a GG-Dh2CpmFc(-) domain, and (ii) a second polypeptide chain comprising a GG-
 15 DhCpmFc(+) domain.

In certain embodiments, a tetramer is provided, comprising a dimer of two GG-Dh2CpmFc(-)-GDF15(N3D):GG-Dh2CpmFc(+) heterodimers in which the two first polypeptide chains of each heterodimer are linked via an interchain disulfide bond between their respective GDF15 regions.

20 More particularly, in a specific embodiment, the tetramer comprises:

(a) two GG-Dh2CpmFc(+) domains (one each heterodimer) comprising the sequence of SEQ ID NO:299,

(b) two GG-Dh2CpmFc(-) domains (one each heterodimer) comprising the sequence of SEQ ID NO:206, and

25 (c) two GDF15(N3D) polypeptides (one each heterodimer) comprising the sequence of SEQ ID NO:52.

In a preferred embodiment, the first polypeptide chain comprises the amino acid sequence:

GGPSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVDGVEVH
 30 NAKTKPREEQYNSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPAPIEKT
 ISKAKGQPREPQVYTLPPSREEMTKNQVSLTCLVKGFYPSDIAVEWESNG
 QPENNYDTTPVLDSGDSFFLYSDLTVDKSRWQQGNVFSCSVMHEALHNH
 YTQKSLSLSPGARDGDHCPLGPGRCRLHTVRSLEDLGWADWVLSPREV
 QVTMCIGACPSQFRAANMHAQIKTSLHRLKPDTPAPCCVPASYNPMVLI
 35 QKTDGTGVSLLQTYDDLLAKDCHCI (SEQ ID NO:211) ,

which is encoded by the nucleic acid sequence:

ggtggcccgctcagtccttctcttccccccaaaacccaaggaca
 ccctcatgatctcccgacccctgaggtcacatgctgggtggacgtg
 agccacgaagaccctgaggtcaagttcaactggtacgtggacggcgtgga
 5 ggtgcataatgccaagacaaagccgcgggaggagcagtacaacagcacgt
 accgtgtgggtcagcgtcctcaccgtcctgcaccaggactggctgaatggc
 aaggagtacaagtgaaggtctccaacaaagccctcccagcccccatcga
 gaaaaccatctccaaagccaaagggcagccccgagaaccacaggtgtaca
 ccctgcccccatcccgggaggagatgaccaagaaccagggtcagcctgacc
 10 tgcttgggtcaaaggcttctatcccagcgacatcgccgtggagtgggagag
 caatgggcagccggagaacaactacgacaccacgcctcccgtgctggact
 ccgacggctccttcttctctatagcgacctcaccgtggacaagagcagg
 tggcagcaggggaacgtcttctcatgctccgtgatgcatgaggctctgca
 caaccactacacgcagaagagcctctccctgtctccgggtgcgcgcgacg
 15 gagaccactgtccgctcgggccccgggcttggctgccgtctgcacacggtc
 cgcgcgtcgtggaagacctgggctgggcccattgggtgctgtcgccacg
 ggaggtgcaagtgacctgtgcatcggcgcgtgcccgagccagttccggg
 cggcaaacatgcacgcgcagatcaagacgagcctgcaccgcctgaagccc
 gacacgggtgccagcgcctgtgctgctgcccgccagctacaatcccatggt
 20 gctcattcaaaagaccgacaccggggtgtcgctccagacctatgatgact
 tgttagccaaagactgccactgcata (SEQ ID NO:210) .

In an embodiment employing the VH21 signal sequence, in a preferred embodiment, the first polypeptide chain comprises the amino acid sequence (signal sequence single underlined):

25 MEWSWVFLFFLSVTTGVHSGGPSVFLFPPKPKDTLMISRTPEVTCVVVDV
 SHEDPEVKFNWYVDGVEVHNAKTKPREEQYNSTYRVVSVLTVLHQDWLNG
 KEYKCKVSNKALPAPIEKTISKAKGQPREPQVYTLPPSREEMTKNQVSLT
 CLVKGFYPSDIAVEWESNGQPENNYDTPPVLDSDGSFFLYSDLTVDKSR
 WQQGNVFSCSVMHEALHNHYTQKSLSLSPGARDGDHCPLGPGRCRLHTV
 30 RASLEDLGWADWVLSPREVQVTMCIGACPSQFRAANMHAQIKTSLHRLKP
 DTVFAPCCVPASYNPMVLIQKTDGTGVSLSLQTYDDLLAKDCHCI (SEQ ID NO:247) ,

which is encoded by the nucleic acid sequence (signal sequence underlined):

atggaatggagctgggtctttctcttcttctcctgtcagtaacgactgggtgt
ccactccgggtggcccgctcagtccttctcttccccccaaaacccaaggaca

ccctcatgatctcccgacccctgaggtcacatgctggtggtggacgtg
agccacgaagaccctgaggtcaagttcaactggtacgtggacggcgtgga
ggtgcataatgccaaagacaaagccgcgggagggagcagtacaacagcacgt
accgtgtggtcagcgtcctcacgctcctgcaccaggactggctgaatggc
5 aaggagtacaagtgaaggtctccaacaaagccctcccagcccccatcga
gaaaaccatctccaaagccaaagggcagcccgagaaccacaggtgtaca
ccctgcccccatccccgggaggagatgaccaagaaccaggtcagcctgacc
tgctgtggtcaaaggttctatcccagcgacatcgccgtggagtgggagag
caatgggcagccggagaacaactacgacaccacgcctcccgtgctggact
10 ccgacggctccttcttctctatagcgacctcaccgtggacaagagcagg
tggcagcaggggaacgtcttctcatgctccgtgatgcatgaggctctgca
caaccactacacgcagaagagcctctccctgtctccgggtgcgcgcgacg
gagaccactgtccgctcgggccccgggcttgctgccgtctgcacacggtc
cgcgcgtcgtggaagacctgggctgggcccattgggtgctgtcgccacg
15 ggaggtgcaagtgaccatgtgcatcggcgcgtgcccagccagttccggg
cggcaaacatgcacgcgcagatcaagacgagcctgcaccgcctgaagccc
gacacgggtgccagcgccctgctgctgcccgcagctacaatcccatggt
gtcattcaaaagaccgacacggggtgtcgtccagacctatgatgact
tgttagccaaagactgccactgcata (SEQ ID NO:248).

20 In a preferred embodiment, the second polypeptide chain comprises the amino acid sequence of SEQ ID NO:205, which is encoded by the nucleic acid sequence of SEQ ID NO:209.

In an embodiment employing the VH21 signal sequence, in a preferred embodiment, the second polypeptide chain comprises the amino acid sequence of SEQ ID NO:245, which
25 is encoded by the nucleic acid sequence of SEQ ID NO:246.

As discussed above, a tetramer is provided comprising two polypeptide chains comprising the sequence of SEQ ID NO:211 and two polypeptide chains comprising the sequence of SEQ ID NO:205.

30 *II.H.28 GG-Dh2CpmFc(-)(Y349C)-GDF15(Ndel3):
GG-Dh2CpmFc(+)(S354C)*

The designation “GG-Dh2CpmFc(-)(Y349C)-GDF15(Ndel3):GG-Dh2CpmFc(+)(S354C)” in the instant disclosure refers to a heterodimer, which comprises (i)
a first polypeptide chain comprising GDF15(Ndel3) polypeptide, the N-terminus of which is
35 linked directly to the C-terminus of a GG-Dh2CpmFc(-)(Y349C) domain, and (ii) a second polypeptide chain comprising a GG-Dh2CpmFc(+)(S345C) domain. The cysteine clamp

mutations allow the first and second polypeptide chains to be linked via an interchain disulfide bond between C349 of the first polypeptide chain and C354 of the second polypeptide chain.

- In certain embodiments, a tetramer is provided, comprising a dimer of two GG-Dh2CpmFc(-)(Y349C)-GDF15(Nde13):GG-Dh2CpmFc(+)(S354C) heterodimers in which the two first polypeptide chains of each heterodimer are linked via an interchain disulfide bond between their respective GDF15 regions.

More particularly, in a specific embodiment, the tetramer comprises:

(a) two GG-Dh2CpmFc(+)(S354C) (one each heterodimer) comprising the sequence:

10 GGPSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVDGVEVH
NAKTKPREEQYNSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPAPIEKT
ISKAKGQPREPQVYTLPPCRKEMTKNQVSLTCLVKGFYPSDIAVEWESNG
QPENNYKTTTPVLKSDGSFFLYSKLTVDKSRWQQGNVFSCSVMHEALHNH
YTQKSLSLSPG (SEQ ID NO:300),

- 15 (b) two GG-Dh2CpmFc(-)(Y349C) chains (one each heterodimer) comprising the sequence:

GGPSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVDGVEVH
NAKTKPREEQYNSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPAPIEKT
ISKAKGQPREPQVCTLPSSREEMTKNQVSLTCLVKGFYPSDIAVEWESNG
20 QPENNYDTTPVLDSGSDGSFFLYSDLTVDKSRWQQGNVFSCSVMHEALHNH
YTQKSLSLSPG (SEQ ID NO:213),

and

(c) two GDF15(Nde13) polypeptides (one each heterodimer) comprising the sequence of SEQ ID NO:55.

- 25 In a preferred embodiment, the first polypeptide chain comprises the amino acid sequence:

GGPSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVDGVEVH
NAKTKPREEQYNSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPAPIEKT
ISKAKGQPREPQVCTLPSSREEMTKNQVSLTCLVKGFYPSDIAVEWESNG
30 QPENNYDTTPVLDSGSDGSFFLYSDLTVDKSRWQQGNVFSCSVMHEALHNH
YTQKSLSLSPGGDHCPLGPGRCRLHTVRASLEDLGWADWVLSPREVQVT
MCIGACPSQFRAANMHAQIKTSLHRLKPDTPAPCCVPASYNPMVLIQKT
DTGVSLQTYDDLAKDCHCI (SEQ ID NO:215),

which is encoded by the nucleic acid sequence:

ggtggcccgctcagtccttctcttccccccaaaacccaaggaca
 ccctcatgatctcccgaccctgaggtcacatgctgggtggacgtg
 agccacgaagaccctgaggtcaagttcaactggtacgtggacggcgtgga
 ggtgcataatgccaaagacaaagccgaggagcagtacaacagcagct
 5 accgtgtggtcagcgtcctcaccgtcctgcaccaggactggctgaatggc
 aaggagtacaagtgaaggtctccaacaaagccctcccagcccccatcga
 gaaaaccatctccaaagccaaagggcagccccgagaaccacaggtgtgca
 ccctgcccccatcccgaggagatgaccaagaaccaggtcagcctgacc
 tgcttgggtcaaaggcttctatcccagcgacatcgccgtggagtgggagag
 10 caatgggcagccggagaacaactacgacaccacgcctcccgtgctggact
 ccgacggctccttcttctctatagcgacctcaccgtggacaagagcagg
 tggcagcaggggaacgtcttctcatgctccgtgatgcatgaggctctgca
 caaccactacacgcagaagagcctctccctgtctccgggtggagaccact
 gtccgctcgggcccggcggttgcgtgcctgtgcacacgggtccgcgcgtcg
 15 ctggaagacctgggctgggcccattgggtgctgctgccacgggaggtgca
 agtgaccatgtgcatcggcgcgtgcccagccagttccgggcccgaaca
 tgcaacgcgcagatcaagacgagcctgcaccgcctgaagcccgcacgggtg
 ccagcgccctgtgctgctgcccgcagctacaatcccatgggtgctcattea
 aaagaccgacaccggggtgctgctccagacctatgatgacttggttagcca
 20 aagactgccactgcata (SEQ ID NO:214).

In an embodiment employing the VH21 signal sequence, in a preferred embodiment, the first polypeptide chain comprises the amino acid sequence (signal sequence single underlined):

MEWSWVFLFFLSVTTGVHSGGPSVFLFPPKPKDTLMISRTPEVTCVVVDV
 25 SHEDPEVKFNWYVDGVEVHNAKTKPREEQYNSTYRVVSVLTVLHQDWLNG
 KEYKCKVSNKALPAPIEKTISKAKGQPREPQVCTLPSSREEMTKNQVSLT
 CLVKGFYPSDIAVEWESNGQPENNYDTPPVLDSDGSFFLYSDLTVDKSR
 WQQGNVFSCSVMHEALHNHYTQKSLSLSPGGDHCP LGPRCCRLHTVRAS
 LEDLGWADWVLSPREVQVTMCIGACPSQFRAANMHAQIKTSLHRLKPDTV
 30 PAPCCVPASYNPMVLIQKTDGTGVS LQTYDDLAKDCHCI (SEQ ID NO:249),

which is encoded by the nucleic acid sequence (signal sequence underlined):

atggaatggagctgggtccttctcttcttctcctgtcagtaacgactgggtgt
 ccactccgggtggcccgtcagtccttcttcttccccccaaaacccaaggaca
 ccctcatgatctcccgaccctgaggtcacatgctgggtggacgtg
 35 agccacgaagaccctgaggtcaagttcaactggtacgtggacggcgtgga

ggtgcataatgccaaagacaaagccgcgggaggagcagtacaacagcacgt
 accgtgtggtcagcgtcctcaccgtcctgcaccaggactggctgaatggc
 aaggagtacaagtgaaggtctccaacaaagccctcccagcccccatcga
 gaaaaccatctccaaagccaaagggcagccccgagaaccacaggtgtgca
 5 cctgcccccatccccggaggagatgaccaagaaccaggtcagcctgacc
 tgctgtggtcaaaggcttctatcccagcgacatcgccgtggagtgggagag
 caatgggcagccggagaacaactacgacaccacgcctcccgtgctggact
 ccgacggctccttcttctctatagcgacctcaccgtggacaagagcagg
 tggcagcaggggaacgtcttctcatgctccgtgatgcatgaggctctgca
 10 caaccactacacgcagaagagcctctccctgtctccgggtggagaccact
 gtccgctcgggccccggcggttgctgccgtctgcacacgggtccgcgcgtcg
 ctggaagacctgggctgggcccattgggtgctgtcgccacgggaggtgca
 agtgaccatgtgcatcggcgcgtgcccgagccagttccgggcggaaca
 tgcacgcgcagatcaagacgagcctgcaccgcctgaagcccagacaggtg
 15 ccagcgcctgctgctgctgccgcccagctacaatcccatggtgctcattca
 aaagaccgacaccggggtgtcgtccagacctatgatgacttgtagcca
 aagactgccactgcata (SEQ ID NO:250) .

In a preferred embodiment, the second polypeptide chain comprises the amino acid sequence:

20 GGPSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVDGVEVH
 NAKTKPREEQYNSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPAPIEKT
 ISKAKGQPREPQVYTLPPCRKEMTKNQVSLTCLVKGFYPSDIAVEWESNG
 QPENNYKTPPVLKSDGSFFLYSKLTVDKSRWQQGNVFSCSVMHEALHNH
 YTQKSLSLSPGK (SEQ ID NO:212) ,

25 which is encoded by the nucleic acid sequence:

ggtggcccgtcagttcttcttccccccaaaaccaaggaca
 cctcatgatctcccgaccctgaggtcacatgctggtggtggacgtg
 agccacgaagacctgaggtcaagttcaactggtacgtggacggcgtgga
 ggtgcataatgccaaagacaaagccgcgggaggagcagtacaacagcacgt
 30 accgtgtggtcagcgtcctcaccgtcctgcaccaggactggctgaatggc
 aaggagtacaagtgaaggtctccaacaaagccctcccagcccccatcga
 gaaaaccatctccaaagccaaagggcagccccgagaaccacaggtgtaca
 ccctgcccccatgccggaaggagatgaccaagaaccaggtcagcctgacc
 tgctgtggtcaaaggcttctatcccagcgacatcgccgtggagtgggagag
 35 caatgggcagccggagaacaactacaagaccacgcctcccgtgctgaagt

ccgacggctccttcttctctatagcaagctcacctggacaagagcagg
 tggcagcaggggaacgtcttctcatgctccgtgatgcatgaggctctgca
 caaccactacacgcagaagagcctctccctgtctccgggtaaa (SEQ ID NO:216) .

- In an embodiment employing the VH21 signal sequence, in a preferred embodiment,
 5 the second polypeptide chain comprises the amino acid sequence (signal sequence single underlined):

MEWSWVFLFFLSVTTGVHSGGPSVFLFPPKPKDTLMISRTPEVTCVVVDV
 SHEDPEVKFNWYVDGVEVHNAKTKPREEQYNSTYRVVSVLTVLHQDWLNG
 KEYKCKVSNKALPAPIEKTISKAKGQPREPQVYTLPPCRKEMTKNQVSLT
 10 CLVKGFYPSDIAVEWESNGQPENNYKTTTPVLKSDGSFFLYSKLTVDKSR
 WQQGNVFSCSVMHEALHNHYTQKSLSLSPGK (SEQ ID NO:251) ,

which is encoded by the nucleic acid sequence (signal sequence underlined):

atggaatggagctgggtcttctcttcttctcctgtcagtaacgactgggtgt
ccactccgggtggcccgctcagtccttcttctcccccaaaacccaaggaca
 15 ccctcatgatctcccgacccctgaggtcacatgctgggtggaggcgtg
 agccacgaagaccctgaggtcaagttcaactggtacgtggacggcgtgga
 ggtgcataatgccaagacaaagccgcgggaggagcagtacaacagcacgt
 accgtgtgggtcagcgtcctcacctgaccaggactgggtgaatggc
 aaggagtacaagtgaaggtctccaacaaagccctcccagccccatcga
 20 gaaaaccatctccaaagccaaagggcagccccgagaaccacaggtgtaca
 ccctgcccccatgccggaaggagatgaccaagaaccagggtcagcctgacc
 tgcttgggtcaaaggcttctatcccagcgacatcgccgtggagtgggagag
 caatgggcagccggagaacaactacaagaccacgcctcccggtgctgaagt
 ccgacggctccttcttctctatagcaagctcacctggacaagagcagg
 25 tggcagcaggggaacgtcttctcatgctccgtgatgcatgaggctctgca
 caaccactacacgcagaagagcctctccctgtctccgggtaaa (SEQ ID NO:252) .

As discussed above, a tetramer is provided comprising two polypeptide chains
 comprising the sequence of SEQ ID NO:215 and two polypeptide chains comprising the
 sequence of SEQ ID NO:212.

30

II.H.29 GG-Dh2CpmFc(-)(Y349C)-GDF15(N3D):GG-Dh2CpmFc(+)(S354C)

- The designation “GG-Dh2CpmFc(-)(Y349C)-GDF15(N3D):GG-
 Dh2CpmFc(+)(S354C)” in the instant disclosure refers to a heterodimer, which comprises (i)
 35 a first polypeptide chain comprising a GDF15(N3D) polypeptide, the N-terminus of which is

linked directly to the C-terminus of a GG-Dh2CpmFc(-)(Y349C) domain, and (ii) a second polypeptide chain comprising a GG-Dh2CpmFc(+)(S345C) domain. The cysteine clamp mutations allow the first and second polypeptide chains to be linked via an interchain disulfide bond between C349 of the first polypeptide chain and C354 of the second polypeptide chain.

In certain embodiments, a tetramer is provided, comprising a dimer of two GG-Dh2CpmFc(-)(Y349C)-GDF15(N3D):GG-Dh2CpmFc(+)(S354C) heterodimers in which the two first polypeptide chains of each heterodimer are linked via an interchain disulfide bond between their respective GDF15 regions.

More particularly, in a specific embodiment, the tetramer comprises:

(a) two GG-Dh2CpmFc(+)(S354C) domains (one each heterodimer) comprising the sequence of SEQ ID NO:300,

(b) two GG-Dh2CpmFc(-)(Y349C) domains (one each heterodimer) comprising the sequence of SEQ ID NO:213, and

(c) two GDF15(N3D) polypeptides (one each heterodimer) comprising the sequence of SEQ ID NO:52.

In a preferred embodiment, the first polypeptide chain comprises the amino acid sequence:

GGPSVFLFPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVDGVEVH
 NAKTKPREEQYNSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPAPIEKT
 ISKAKGQPREPQVCTLPPSREEMTKNQVSLTCLVKGFYPSDIAVEWESNG
 QPENNYDTTPVLDSGDSFFLYSDLTVDKSRWQQGNVFSCSVMHEALHNH
 YTQKSLSLSPGARDGDHCPLGPGRCRLHTVRASLEDLGWADWVLSPREV
 QVTMCIGACPSQFRAANMHAQIKTSLHRLKPDTPAPCCVPASYNPMVLI
 QKTDGTGVS LQTYDDL LAKDCHCI (SEQ ID NO:218),

which is encoded by the nucleic acid sequence:

ggtggcccgctcagtccttcctcttccccccaaaacccaaggaca
 ccctcatgatctcccgaccctgaggtcacatgctgggtggacgtg
 agccacgaagaccctgaggtcaagttcaactggtacgtggacggcgtgga
 ggtgcataatgccaaagacaaagccgcgggaggagcagtacaacagcacgt
 accgtgtggtcagcgtcctcaccgtcctgcaccaggactggctgaatggc
 aaggagtacaagtgaaggtctccaacaaagccctccagcccccatcga
 gaaaaccatctccaaagccaaagggcagccccgagaaccacaggtgtgca
 ccctgcccccatcccgaggagatgaccaagaaccaggtcagcctgacc
 tgccctggtcaaaggcttctatcccagcgacatcgccgtggagtgggagag

caatgggcagccggagaacaactacgacaccacgcctcccgtgctggact
 ccgacggctccttcttctctatagcgacctcaccgtggacaagagcagg
 tggcagcaggggaacgtcttctcatgctccgtgatgcatgaggctctgca
 caaccactacacgcagaagagcctctccctgtctccgggtgcgcgcgaag
 5 gagaccactgtccgctcgggccccggggttgcgtgccgtctgcacacggtc
 cgcgcgtcgtggaagacctgggctgggcccattgggtgctgtcgccacg
 ggagggtgcaagtgacctgtgcatcggcgcgtgcccagccagttccggg
 cggcaaacatgcacgcgcagatcaagacgagcctgcaccgcctgaagccc
 gacacgggtgccagcgcctgtgctgctgcccgcagctacaatcccatggt
 10 gctcattcaaaagaccgacaccggggtgtcgtcctcagacctatgatgact
 tgttagccaaagactgccactgcata (SEQ ID NO:217) .

In an embodiment employing the VH21 signal sequence, in a preferred embodiment, the first polypeptide chain comprises the amino acid sequence (signal sequence single underlined):

15 MEWSWVFLFFLSVTTGVHSGGPSVFLFPPKPKDTLMISRTPEVTCVVVDV
 SHEDPEVKFNWYVDGVEVHNAKTKPREEQYNSTYRVVSVLTVLHQDWLNG
 KEYKCKVSNKALPAPIEKTISKAKGQPREPQVCTLPSSREEMTKNQVSLT
 CLVKGFYPSDIAVEWESNGQPENNYDTTPVLDSGDSFFLYSDLTVDKSR
 WQQGNVFCFSVMHEALHNHYTQKSLSLSPGARDGHCPLGPGRCCRLHTV
 20 RASLEDLGWADWVLSPREVQVTMCIGACPSQFRAANMHAQIKTSLHRLKP
 DTVPAAPCCVPASYNPMVLIQKTDGTGVS LQTYDDLAKDCHCI (SEQ ID NO:253) ,

which is encoded by the nucleic acid sequence (signal sequence underlined):

atggaatggagctgggtctttctcttcttctcctgtcagtaacgactgggtgt
ccactccgggtggcccgtcagtccttcttctcccccaaaacccaaggaca
 25 cctcatgatctcccgaccctgaggtcacatgcgtgggtggacgtg
 agccacgaagacctgaggtcaagttcaactggtacgtggacggcgtgga
 ggtgcataatgccaaagacaaagccgcgggaggagcagtacaacagcacgt
 accgtgtggtcagcgtcctcaccgtcctgcaccaggactggctgaatggc
 aaggagtacaagtgaaggtctccaacaaagccctcccagcccccatcga
 30 gaaaaccatctccaaagccaaagggcagccccgagaaccacaggtgtgca
 ccctgcccccatcccgggaggagatgaccaagaaccaggtcagcctgacc
 tgccctgggtcaaaggcttctatcccagcgacatcgccgtggagtgggagag
 caatgggcagccggagaacaactacgacaccacgcctcccgtgctggact
 ccgacggctccttcttctctatagcgacctcaccgtggacaagagcagg
 35 tggcagcaggggaacgtcttctcatgctccgtgatgcatgaggctctgca

caaccactacacgcagaagagcctctccctgtctccgggtgcgcgcgacg
 gagaccactgtccgctcgggccccggcggttgctgccgtctgcacacggtc
 cgcgcgtcgttggaagacctgggctgggcccattgggtgctgtcgccacg
 ggaggtgcaagtgaccatgtgcacatggcgctgcccagaccagttccggg
 5 cggcaaacatgcacgcgcagatcaagacgagcctgcaccgcctgaagccc
 gacacgggtgccagcgcctgtgctgctgcccgccagctacaatcccatggt
 gctcattcaaaagaccgacaccggggtgtcgtccagacctatgatgact
 tgttagccaaagactgccactgcata (SEQ ID NO:254).

In a preferred embodiment, the second polypeptide chain comprises the amino acid
 10 sequence of SEQ ID NO:212, which is encoded by the nucleic acid sequence of SEQ ID
 NO:218.

In an embodiment employing the VH21 signal sequence, in a preferred embodiment,
 the second polypeptide chain comprises the amino acid sequence of SEQ ID NO:251, which
 is encoded by the nucleic acid sequence of SEQ ID NO:252.

15 As discussed above, a tetramer is provided comprising two polypeptide chains
 comprising the sequence of SEQ ID NO:218 and two polypeptide chains comprising the
 sequence of SEQ ID NO:212.

II.H.30 Dh3CpmFc(-)-GDF15(Ndel3):Dh3CpmFc(+)

20 The designation “Dh3CpmFc(-)-GDF15(Ndel3):Dh3CpmFc(+)” in the instant
 disclosure refers to a heterodimer, which comprises (i) a first polypeptide chain comprising a
 GDF15(Ndel3) polypeptide, the N-terminus of which is linked directly to the C-terminus of
 a Dh3CpmFc(-) domain, and (ii) a second polypeptide chain comprising a Dh3CpmFc(+)
 domain.

25 In certain embodiments, a tetramer is provided, comprising a dimer of two
 Dh3CpmFc(-)-GDF15(Ndel3):Dh3CpmFc(+) heterodimers in which the two first polypeptide
 chains of each heterodimer are linked via an interchain disulfide bond between their
 respective GDF15 regions.

More particularly, in a specific embodiment, the tetramer comprises:

30 (a) two Dh3CpmFc(+) domains (one each heterodimer) comprising the sequence:

GPSVFLFPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVDGVEVHN
 AKTKPREEQYNSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPAPIEKTI
 SKAKGQPREPQVYTLPPSRKEMTKNQVSLTCLVKGFYPSDIAVEWESNGQ
 PENNYKTPPVLKSDGSFFLYSKLTVDKSRWQQGNVFSVSMHEALHNHY
 35 TQKSLSLSPG (SEQ ID NO:301),

(b) two Dh3CpmFc(-) domains (one each heterodimer) comprising the sequence:

GPSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVDGVEVHN
AKTKPREEQYNSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPAPIEKT I
SKAKGQPREPQVYTLPPSREEMTKNQVSLTCLVKGFYPSDIAVEWESNGQ
5 PENNYDTTPVLDSGDSFFLYSDLTVDKSRWQQGNVFSCSVMHEALHNHY
TQKSLSLSPG (SEQ ID NO:220),

and

(c) two GDF15(Nde13) polypeptides (one each heterodimer) comprising the sequence of SEQ ID NO:55.

10 In a preferred embodiment, the first polypeptide chain comprises the amino acid sequence:

GPSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVDGVEVHN
AKTKPREEQYNSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPAPIEKT I
SKAKGQPREPQVYTLPPSREEMTKNQVSLTCLVKGFYPSDIAVEWESNGQ
15 PENNYDTTPVLDSGDSFFLYSDLTVDKSRWQQGNVFSCSVMHEALHNHY
TQKSLSLSPGGDHCPGGRCCRLHTVRASLEDLGWADWVLSPREVQVTM
CIGACPSQFRAANMHAQIKTSLHRLKPDTPAPCCVPASYNPMVLIQKTD
TGVSLQTYDDLLAKDCHCI (SEQ ID NO:222),

which is encoded by the nucleic acid sequence:

20 ggcccgctcagttcttctcttccccccaaaaccca
aggacaccctcatgatctcccgaccctgaggtcacatgcgtggtggtg
gacgtgagccacgaagaccctgaggtcaagttcaactggtacgtggacgg
cgtggaggtgcataatgccaaagacaaagccgcgggaggagcagtacaaca
gcacgtaccgtgtggtcagcgtcctcaccgtcctgcaccaggactggctg
25 aatggcaaggagtacaagtgaaggtctccaacaaagccctcccagcccc
catcgagaaaaccatctccaaagccaaagggcagccccgagaaccacagg
tgtacaccctgcccccatcccgaggagatgaccaagaaccagggtcagc
ctgacctgcctggtcaaaggcttctatcccagcgacatcgccgtggagtg
ggagagcaatgggcagccggagaacaactacgacaccacgcctcccgtgc
30 tggactccgacggctccttcttctctatagcgacctcaccgtggacaag
agcagggtggcagcaggggaacgtcttctcatgctccgtgatgcatgaggc
tctgcacaaccactacacgcagaagagcctctccctgtctccgggtggag
accactgtccgctcgggccccggcggttgcgtgccgtctgcacacgggtccgc
gcgtcgctggaagacctgggctgggcccattgggtgctgtcgccacggga

ggtgcaagtgaccatgtgcatcggcgcgtgcccagagccagttccgggcg
 caaacatgcacgcgcagatcaagacgagcctgcaccgcctgaagcccgac
 acggtgccagcgccctgctgctgctgcccgcagctacaatcccatggtgct
 cattcaaaagaccgacacccgggtgtcgtccagacctatgatgacttgt
 5 tagccaaagactgccactgcata (SEQ ID NO:221).

In an embodiment employing the VK1 signal sequence, in a preferred embodiment, the first polypeptide chain comprises the amino acid sequence (signal sequence single underlined):

MDMRVPAQLLGLLLLWLRGARGGPSVFLFPPKPKDTLMISRTPEVTCVVV
 10 DVSHEDPEVKFNWYVDGVEVHNAKTKPREEQYNSTYRVVSVLTVLHQDWL
 NGKEYKCKVSNKALPAPIEKTISKAKGQPREPQVYTLPPSREEMTKNQVS
 LTCLVKGFYPSDIAVEWESNGQPENNYDTTPVLDSGSSFFLYSDLTVDK
 SRWQQGNVFSQSVMEALHNHYTQKSLSLSPGGDHCPLGPGRCRLHTVR
 ASLEDLGWADWVLSPREVQVTMCIGACPSQFRAANMHAQIKTSLHRLKPD
 15 TVPAPCCVPASYNPMVLIQKTDGTGVSLQTYDDLAKDCHCI (SEQ ID NO:255),

which is encoded by the nucleic acid sequence (signal sequence underlined):

atggacatgaggggtgcccgtcagctcctggggctcctgctgctgtggct
gagaggtgcgcgctgtggcccgtcagtccttcctcttccccccaaaaccca
 aggacaccctcatgatctcccgaccctgaggtcacatgcgtggtggtg
 20 gacgtgagccacgaagaccctgaggtcaagttcaactggtacgtggacgg
 cgtggaggtgcataatgccaagacaaagccgcgggaggagcagtacaaca
 gcacgtaccgtgtggtcagcgtcctcaccgtcctgcaccaggactggctg
 aatggcaaggagtacaagtgcaaggtctccaacaaagccctcccagcccc
 catcgagaaaaccatctccaaagccaaagggcagccccgagaaccacagg
 25 tgtacaccctgcccccatccgggaggagatgaccaagaaccaggtcagc
 ctgacctgcctggtcaaaggcttctatcccagcgacatcgccgtggagtg
 ggagagcaatgggcagccggagaacaactacgacaccacgcctcccgtgc
 tggactccgacggctccttcttctctatagcgacctcaccgtggacaag
 agcaggtggcagcaggggaacgtcttctcatgctccgtgatgcatgagggc
 30 tctgcacaaccactacacgcagaagagcctctccctgtctccgggtggag
 accactgtccgctcgggccccggcggttgcgtgccgtctgcacacggctccgc
 gcgtcgctggaagacctgggctgggcccattgggtgctgtcgccacggga
 ggtgcaagtgaccatgtgcatcggcgcgtgcccagagccagttccgggcg
 caaacatgcacgcgcagatcaagacgagcctgcaccgcctgaagcccgac
 35 acggtgccagcgccctgctgctgctgcccgcagctacaatcccatggtgct

cattcaaaagaccgacacccgggtgtcgctccagacctatgatgacttgt
tagccaaagactgccactgcata (SEQ ID NO:256) .

In a preferred embodiment, the second polypeptide chain comprises the amino acid sequence:

5 GPSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVDGVEVHN
AKTKPREEQYNSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPAPIEKT
SKAKGQPREPQVYTLPPSRKEMTKNQVSLTCLVKGFYPSDIAVEWESNGQ
PENNYKTTTPVLKSDGSFFLYSKLTVDKSRWQQGNVSCSVMHEALHNHY
TQKSLSLSPGK (SEQ ID NO:219) ,

10 which is encoded by the nucleic acid sequence:

ggcccgctcagtcttcctcttccccccaaaaccca
aggacaccctcatgatctcccgaccctgaggtcacatgcgtggtggtg
gacgtgagccacgaagaccctgaggtcaagttcaactggtacgtggacgg
cgtggaggtgcataatgccaaagacaaagccgcgggaggagcagtacaaca
15 gcacgtaccgtgtggtcagcgtcctcaccgtcctgcaccaggactggctg
aatggcaaggagtacaagtgaaggtctccaacaaagccctcccagcccc
catcgagaaaaccatctccaaagccaaagggcagccccgagaaccacagg
tgtacaccctgcccccatcccggaaggagatgaccaagaaccaggtcagc
ctgacctgcctggtcaaaggcttctatcccagcgacatcgccgtggagtg
20 ggagagcaatgggcagccggagaacaactacaagaccacgcctcccgtgc
tgaagtccgacgggtccttcttctctatagcaagctcacctggacaag
agcaggtggcagcaggggaacgtcttctcatgctccgtgatgcatgaggc
tctgcacaaccactacacgcagaagagcctctccctgtctccgggtaaa
(SEQ ID NO:223) .

25 In an embodiment employing the VK1 signal sequence, in a preferred embodiment,
the second polypeptide chain comprises the amino acid sequence (signal sequence single
underlined):

MDMRVPAQLLGLLLLWLRGARCGPSVFLFPPKPKDTLMISRTPEVTCVVV
DVSHEDPEVKFNWYVDGVEVHNAKTKPREEQYNSTYRVVSVLTVLHQDWL
30 NGKEYKCKVSNKALPAPIEKTISKAKGQPREPQVYTLPPSRKEMTKNQVS
LTCLVKGFYPSDIAVEWESNGQPENNYKTTTPVLKSDGSFFLYSKLTVDK
SRWQQGNVSCSVMHEALHNHYTQKSLSLSPGK (SEQ ID NO:257) ,

which is encoded by the nucleic acid sequence (signal sequence underlined):

atggacatgaggggtgcccgtcagctcctggggctcctgctgctgtggct
 gagagggtgcgcgctgtggcccgtcagtccttctcttccccccaaaaccca
 aggacaccctcatgatctcccgaccctgaggtcacatgcgtgggtggg
 gacgtgagccacgaagaccctgaggtcaagttcaactggtacgtggacgg
 5 cgtggaggtgcataatgccaaagacaaagccgcgggaggagcagtacaaca
 gcacgtaccgtgtggtcagcgtcctcaccgtcctgcaccaggactggctg
 aatggcaaggagtacaagtgaaggtctccaacaaagccctcccagcccc
 catcgagaaaaccatctccaaagccaaagggcagccccgagaaccacagg
 tgtacaccctgcccccatcccgaaggagatgaccaagaaccagggtcagc
 10 ctgacctgcctggtcaaaggcttctatcccagcgacatcgccgtggagtg
 ggagagcaatgggcagccggagaacaactacaagaccacgcctcccgtgc
 tgaagtccgacgggtccttcttctctatagcaagctcaccgtggacaag
 agcaggtggcagcaggggaacgtcttctcatgctccgtgatgcatgaggc
 tctgcacaaccactacacgcagaagagcctctccctgtctccgggtaaa (SEQ ID
 15 NO:258) .

As discussed above, a tetramer is provided comprising two polypeptide chains comprising the sequence of SEQ ID NO:222 and two polypeptide chains comprising the sequence of SEQ ID NO:219.

20 *II.H.31 Dh3CpmFc(-)-GDF15(N3D):Dh3CpmFc(+)*

The designation “Dh3CpmFc(-)-GDF15(N3D):Dh3CpmFc(+)” in the instant disclosure refers to a heterodimer comprising (i) a first polypeptide chain comprising a GDF15(N3D) polypeptide, the N-terminus of which is linked directly to the C-terminus of a Dh3CpmFc(-) domain, and (ii) a second polypeptide chain comprising a Dh3CpmFc(+) domain.
 25

In certain embodiments, a tetramer is provided, comprising a dimer of two Dh3CpmFc(-)-GDF15(N3D):Dh3CpmFc(+) heterodimers in which the two first polypeptide chains of each heterodimer are linked via an interchain disulfide bond between their respective GDF15 regions.

30 More particularly, in a specific embodiment, the tetramer comprises:

(a) two Dh3CpmFc(+) domains (one each heterodimer) comprising the sequence of SEQ ID NO:301,

(b) two Dh3CpmFc(-) (one each heterodimer) comprising the sequence of SEQ ID NO:220, and

35 (c) two GDF15(N3D) polypeptides (one each heterodimer) comprising the sequence of SEQ ID NO:52.

In a preferred embodiment, the first polypeptide chain comprises the amino acid sequence:

GPSVFLFPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVDGVEVHN
AKTKPREEQYNSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPAPIEKTI
5 SKAKGQPREPQVYTLPPSREEMTKNQVSLTCLVKGFYPSDIAVEWESNGQ
PENNYDTPPVLDSDGSFFLYSDLTVDKSRWQQGNVFSCSVMHEALHNHY
TQKSLSLSPGARDGDHCPLGPGRCCRLHTVRASLEDLGWADWVLSPREVQ
VTMCIGACPSQFRAANMHAQIKTSLHRLKPDTPAPCCVPASYNPMVLIQ
KTDGTGVS LQTYDDLAKDCHCI (SEQ ID NO:225),

10 which is encoded by the nucleic acid sequence:

atggacatgagggcgcccgctcagctcctggggctcctgctgctgtggct
gagaggcgcgctgtggcccgctcagtcctcctcttccccccaaaaccca
aggacaccctcatgatctcccgaccctgaggtcacatgcgtgggtggg
gacgtgagccacgaagaccctgaggtcaagttcaactggtacgtggacgg
15 cgtggaggtgcataatgccaaagacaaagccgaggaggagcagtacaaca
gcacgtaccgtgtggtcagcgctcctcaccgtcctgcaccaggactggctg
aatggcaaggagtacaagtgaaggtctccaacaaagccctcccagcccc
catcgagaaaaccatctccaaagccaaagggcagccccgagaaccacagg
tgtacaccctgcccccatcccgaggagatgaccaagaaccaggctcagc
20 ctgacctgcctggtcaaaggcttctatcccagcgacatcgccgtggagtg
ggagagcaatgggcagccggagaacaactacgacaccacgcctcccgtgc
tggactccgacggctccttcttctctatagcgacctcaccgtggacaag
agcagggtggcagcaggggaacgtcttctcatgctccgtgatgcatgaggc
tctgcacaaccactacacgcagaagagcctctccctgtctccgggtgcgc
25 gcgacggagaccactgtccgctcgggcccgggcgttgctgccgtctgcac
acgggtccgcgcgtcgctggaagacctgggctgggcccattgggtgctgtc
gccacgggaggtgcaagtgacctgtgcatcggcgcgtgccgagccagt
tccggggcggaacatgcacgcgcagatcaagacgagcctgcaccgcctg
aagcccgcacgggtgccagcgccctgctgcgtgcccgcagctacaatcc
30 catgggtgctcattcaaaagaccgacaccgggtgtcgctccagacctatg
atgacttgtagccaaagactgccactgcata (SEQ ID NO:224).

In an embodiment employing the VK1 signal sequence, in a preferred embodiment, the first polypeptide chain comprises the amino acid sequence (signal sequence single underlined):

MDMRVPAQLLGLLLLWLRGARGGPSVFLFPPKPKDTLMISRTPEVTCVVV
 DVSHEDPEVKFNWYVDGVEVHNAKTKPREEQYNSTYRVVSVLTVLHQDWL
 NGKEYKCKVSNKALPAPIEKTISKAKGQPREPQVYTLPPSREEMTKNQVS
 LTCLVKGFYPSDIAVEWESNGQPENNYDTTPFVLDSGDSFFLYSDLTVDK
 5 SRWQQGNVFSQSVMEALHNHYTQKSLSLSPGARDGDHCPLGPGRCRLH
 TVRASLEDLGWADWVLSPREVQVTMCIGACPSQFRAANMHAQIKTSLHRL
 KPDTVPAPCCVPASYNPMVLIQKTDGTGVSLLQTYDDLAKDCHCI (SEQ ID NO:259) ,

which is encoded by the nucleic acid sequence (signal sequence underlined):

atggacatgaggggtgcccgctcagctcctggggctcctgctgctgtggct
 10 gagaggtgcgcgctgtggcccgctcagtccttcctcttccccccaaaaccca
 aggacaccctcatgatctcccgaccctgaggtcacatgctgtggtggtg
 gacgtgagccacgaagaccctgaggtcaagttcaactggtacgtggacgg
 cgtggaggtgcataatgccaaagacaaagccgaggaggagcagtacaaca
 gcacgtaccgtgtggtcagcgtcctcaccgtcctgcaccaggactggctg
 15 aatggcaaggagtacaagtgcaaggtctccaacaaagccctcccagcccc
 catcgagaaaaccatctccaaagccaaagggcagccccgagaaccacagg
 tgtacaccctgcccccatcccgaggagatgaccaagaaccagggtcagc
 ctgacctgcctggtcaaaggcttctatcccagcgacatcgccgtggagtg
 ggagagcaatgggcagccggagaacaactacgacaccacgcctcccgtgc
 20 tggactccgacggctccttcttctctatagcgacctcaccgtggacaag
 agcaggtggcagcaggggaacgtcttctcatgctccgtgatgcatgaggc
 tctgcacaaccactacacgcagaagagcctctccctgtctccgggtgcgc
 gcgacggagaccactgtccgctcgggcccggcggttgcctgctgcctgac
 acggtccgcgcgtcgctggaagacctgggctgggccgattgggtgctgtc
 25 gccacgggaggtgcaagtgacctgtgcatcggcgctgcccagaccagt
 tccgggcggaacatgcacgcgcagatcaagacgagcctgcaccgcctg
 aagcccgacacggtgccagcgccctgctgctgcccgcagctacaatcc
 catgggtgctcattcaaaagaccgacaccggggtgtcgctccagacctatg
 atgacttggttagccaaagactgccactgcata (SEQ ID NO:260) .

30 In a preferred embodiment, the second polypeptide chain comprises the amino acid sequence of SEQ ID NO:219, which is encoded by the nucleic acid sequence of SEQ ID NO:223.

In an embodiment employing the VK1 signal sequence, in a preferred embodiment, the second polypeptide chain comprises the amino acid sequence of SEQ ID NO:257, which
 35 is encoded by the nucleic acid sequence of SEQ ID NO:258.

As discussed above, a tetramer is provided comprising two polypeptide chains comprising the sequence of SEQ ID NO:225 and two polypeptide chains comprising the sequence of SEQ ID NO:219.

5 *II.H.32 Dh3CpmFc(-)(Y349C)-GDF15(Ndel3):Dh3CpmFc(+)(S354C)*

The designation “Dh3CpmFc(-)(Y349C)-GDF15(Ndel3):Dh3CpmFc(+)(S354C)” in the instant disclosure refers to a heterodimer, which comprises (i) a first polypeptide chain comprising a GDF15(Ndel3) polypeptide, the N-terminus of which is linked directly to the C-terminus of a Dh3CpmFc(-)(Y349C) domain, and (ii) a second polypeptide chain comprising
10 a Dh3CpmFc(+)(S345C) domain. The cysteine clamp mutations allow the first and second polypeptide chains to be linked via an interchain disulfide bond between C349 of the first polypeptide and C354 of the second polypeptide.

In certain embodiments, a tetramer is provided, comprising a dimer of two Dh3CpmFc(-)(Y349C)-GDF15(Ndel3):Dh3CpmFc(+)(S354C) heterodimers in which the two
15 first polypeptide chains of each heterodimer are linked via an interchain disulfide bond between their respective GDF15 regions.

More particularly, in a specific embodiment, the tetramer comprises:

(a) two Dh3CpmFc(+)(S354C) domains (one each heterodimer) comprising the sequence:

20 GPSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVDGVEVHN
AKTKPREEQYNSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPAPIEKT
SKAKGQPREPQVYTLPPCRKEMTKNQVSLTCLVKGFYPSDIAVEWESNGQ
PENNYKTPPVLKSDGSFFLYSKLTVDKSRWQQGNVFSVMSVHEALHNHY
TQKSLSLSPG (SEQ ID NO:302),

25 (b) two Dh3CpmFc(-)(Y349C) domains (one each heterodimer) comprising the sequence:

GPSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVDGVEVHN
AKTKPREEQYNSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPAPIEKT
SKAKGQPREPQVCTLPSSREEMTKNQVSLTCLVKGFYPSDIAVEWESNGQ
30 PENNYDTTPPVLDSDGSFFLYSDLTVDKSRWQQGNVFSVMSVHEALHNHY
TQKSLSLSPG (SEQ ID NO:227),

and

(c) two GDF15(Ndel3) polypeptides (one each heterodimer) comprising the sequence of SEQ ID NO:55.

In a preferred embodiment, the first polypeptide chain comprises the amino acid sequence:

GPSVFLFPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVDGVEVHN
AKTKPREEQYNSTYRVSVLTVTLHQDWLNGKEYKCKVSNKALPAPIEKTI
5 SKAKGQPREPQVCTLPSSREEMTKNQVSLTCLVKGFYPSDIAVEWESNGQ
PENNYDTPPVLDSDGSFFLYSDLTVDKSRWQQGNVFSCSVMHEALHNHY
TQKSLSLSPGGDHCPLGPGRCCRLHTVRASLEDLGWADWVLSPREVQVTM
CIGACPSQFRAANMHAQIKTSLHRLKPDTPAPCCVPASYNPMVLIQKTD
TGVSLQTYDDLAKDCHCI (SEQ ID NO:229),

10 which is encoded by the nucleic acid sequence:

atggacatgagggcgcccgctcagctcctggggctcctgctgctgtggct
gagaggtgcgcgctgtggcccgctcagtcctcctcttccccccaaaaccca
aggacaccctcatgatctcccgaccctgaggtcacatgcgtgggtggg
gacgtgagccacgaagaccctgaggtcaagttcaactggtacgtggacgg
15 cgtggaggtgcataatgccaaagacaaagccgcgaggagcagtacaaca
gcacgtaccgtgtggtcagcgtcctcaccgtcctgcaccaggactggctg
aatggcaaggagtacaagtgaaggtctccaacaaagccctcccagcccc
catcgagaaaaccatctccaaagccaaagggcagccccgagaaccacagg
tgtgcaccctgcccccatcccgaggagatgaccaagaaccagggtcagc
20 ctgacctgcctggtcaaaggcttctatcccagcgacatcgccgtggagtg
ggagagcaatgggcagccggagaacaactacgacaccacgcctcccgtgc
tggactccgacgggtccttcttctctatagcgacctcaccgtggacaag
agcaggtggcagcaggggaacgtcttctcatgctccgtgatgcatgaggc
tctgcacaaccactacacgcagaagagcctctccctgtctccgggtggag
25 accactgtccgctcgggccccggcggttgetgccgtctgcacacgggtccgc
gcgtcgctggaagacctgggctgggcccattgggtgctgtcgccacggga
gggtgcaagtgacctgtgcatcggcgcgtgcccagccagttccgggcgg
caaacatgcacgcgcagatcaagacgagcctgcaccgcctgaagcccgac
acgggtgccagcgccctgctgctgcccgcagctacaatcccatggtgct
30 cattcaaaagaccgacaccgggggtgctgctccagacctatgatgacttgt
tagccaaagactgccactgcata (SEQ ID NO:228).

In an embodiment employing the VK1 signal sequence, in a preferred embodiment, the first polypeptide chain comprises the amino acid sequence (signal sequence single underlined):

MDMRVPAQLLGLLLLWLRGARGGPSVFLFPPKPKDTLMISRTPEVTCVVV
 DVSHEDPEVKFNWYVDGVEVHNAKTKPREEQYNSTYRVVSVLTVLHQDWL
 NGKEYKCKVSNKALPAPIEKTISKAKGQPREPQVCTLPSPREEMTKNQVS
 LTCLVKGFYPSDIAVEWESNGQPENNYDTTPVLDSGSSFFLYSDLTVDK
 5 SRWQQGNVFSQSVMEALHNHYTQKSLSLSPGGDHCPLGPRCCRLHTR
 ASLEDLGWADWVLSPREVQVTMCIGACPSQFRAANMHAQIKTSLHRLKPD
 TVPAPCCVPASYNPMVLIQKTDGTGVSQTYDDLLAKDCHCI (SEQ ID NO:261),

which is encoded by the nucleic acid sequence (signal sequence underlined):

atggacatgaggggtgcccgctcagctcctggggctcctgctgctgtggct
 10 gagaggtgcgcgctgtggcccgctcagtccttcttccccccaaaaccca
 aggacaccctcatgatctcccgaccctgaggtcacatgctggtggtg
 gacgtgagccacgaagaccctgaggtcaagttcaactggtacgtggacgg
 cgtggaggtgcataatgccaaagacaaagccgaggagagcagtacaaca
 gcacgtaccgtgtggtcagcgtcctcaccgtcctgcaccaggactggctg
 15 aatggcaaggagtacaagtgaaggtctccaacaaagccctcccagcccc
 catcgagaaaaccatctccaaagccaaagggcagccccgagaaccacagg
 tgtgcaccctgcccccatcccgaggagatgaccaagaaccagggtcagc
 ctgacctgcctggtcaaaggcttctatcccagcgacatcgccgtggagtg
 ggagagcaatgggcagccggagaacaactacgacaccacgcctcccgtgc
 20 tggactccgacggctccttcttctctatagcgacctcaccgtggacaag
 agcaggtggcagcaggggaacgtcttctcatgctccgtgatgcatgaggc
 tctgcacaaccactacacgcagaagagcctctccctgtctccgggtggag
 accactgtccgctcgggccccggcggttgcgtgccgtctgcacacggtcgc
 gcgtcgctggaagacctgggctgggcccattgggtgctgctgccacggga
 25 ggtgcaagtgaccatgtgcatcggcgcgtgcccagaccagttccgggagg
 caaacatgcacgcgagatcaagacgagcctgcaccgcctgaagcccgac
 acggtgccagcgccctgctgcgtgcccgcagctacaatcccatggtgct
 cattcaaaagaccgacaccgggtgtcgctccagacctatgatgacttgt
 tagccaaagactgccactgcata (SEQ ID NO:262).

30 In a preferred embodiment, the second polypeptide chain comprises the amino acid sequence:

GPSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVDGVEVHN
 AKTKPREEQYNSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPAPIEKT
 ISKAKGQPREPQVYTLPPCRKEMTKNQVSLTCLVKGFYPSDIAVEWESNGQ

PENNYKTTTPVLKSDGSFFLYSKLTVDKSRWQQGNVFSCSVMHEALHNHY
TQKSLSLSPGK (SEQ ID NO: 226),

which is encoded by the nucleic acid sequence:

ggcccgctcagtccttcctcttccccccaaaaccca
5 aggacaccctcatgatctcccggaccctgaggtcacatgcgtggtggtg
gacgtgagccacgaagaccctgaggtcaagttcaactggtacgtggacgg
cgtggaggtgcataatgccaaagacaaagccgcgggaggagcagtacaaca
gcacgtaccgtgtggtcagcgtcctcaccgtcctgcaccaggactggctg
aatggcaaggagtacaagtgaaggtctccaacaaagccctcccagcccc
10 catcgagaaaaccatctccaaagccaaagggcagccccgagaaccacagg
tgtacaccctgcccccatgccggaaggagatgaccaagaaccagggtcagc
ctgacctgcctggtcaaaggcttctatcccagcgacatcgccgtggagtg
ggagagcaatgggcagccggagaacaactacaagaccacgcctcccgtgc
tgaagtccgacggctccttcttctctatagcaagctcaccgtggacaag
15 agcaggtggcagcaggggaacgtcttctcatgctccgtgatgcatgaggc
tctgcacaaccactacacgcagaagagcctctccctgtctccgggtaaa
(SEQ ID NO: 230).

In an embodiment employing the VK1 signal sequence, in a preferred embodiment,
the second polypeptide chain comprises the amino acid sequence (signal sequence single
20 underlined):

MDMRVPAQLLGLLLLWLRGARGPSVFLFPPKPKDTLMISRTPEVTCVVV
DVSHEDPEVKFNWYVDGVEVHNAKTKPREEQYNSTYRVVSVLTVLHQDWL
NGKEYKCKVSNKALPAPIEKTISKAKGQPREPQVYTLPPCRKEMTKNQVS
LTCLVKGFYPSDIAVEWESNGQPENNYKTTTPVLKSDGSFFLYSKLTVDK
25 SRWQQGNVFSCSVMHEALHNHYTQKSLSLSPGK (SEQ ID NO: 263),

which is encoded by the nucleic acid sequence (signal sequence underlined):

atggacatgaggggtgcccgctcagctcctggggctcctgctgctgtggct
gagaggtgcgcgctgtggcccgtcagtccttcctcttccccccaaaaccca
aggacaccctcatgatctcccggaccctgaggtcacatgcgtggtggtg
30 gacgtgagccacgaagaccctgaggtcaagttcaactggtacgtggacgg
cgtggaggtgcataatgccaaagacaaagccgcgggaggagcagtacaaca
gcacgtaccgtgtggtcagcgtcctcaccgtcctgcaccaggactggctg
aatggcaaggagtacaagtgaaggtctccaacaaagccctcccagcccc
catcgagaaaaccatctccaaagccaaagggcagccccgagaaccacagg

5 tgtacaccctgcccccatgccggaaggagatgaccaagaaccagggtcagc
 ctgacctgcctgggtcaaaggcttctatcccagcgacatcgccgtggagtg
 ggagagcaatgggcagccggagaacaactacaagaccacgcctcccgtgc
 tgaagtccgacgggtccttcttctctatagcaagctcacctgggacaag
 agcagggtggcagcaggggaacgtcttctcatgctccgtgatgcatgaggc
 tctgcacaaccactacacgcagaagagcctctccctgtctccgggtaaa (SEQ ID
 NO:264) .

As discussed above, a tetramer is provided comprising two polypeptide chains
 comprising the sequence of SEQ ID NO:229 and two polypeptide chains comprising the
 10 sequence of SEQ ID NO:226.

II.H.33 Dh3CpmFc(-)(Y349C)-GDF15(N3D):Dh3CpmFc(+)(S354C)

The designation “Dh3CpmFc(-)(Y349C)-GDF15(N3D):Dh3CpmFc(+)(S354C)” in
 the instant disclosure refers to a heterodimer, which comprises (i) a first polypeptide chain
 15 comprising a GDF15(N3D) polypeptide, the N-terminus of which is linked directly to the C-
 terminus of a Dh3CpmFc(-)(Y349C) domain, and (ii) a second polypeptide chain comprising
 a Dh3CpmFc(+)(S354C) domain. The cysteine clamp mutations allow the first and second
 polypeptide chains to be linked via an interchain disulfide bond between C349 of the first
 polypeptide chain and C354 of the second polypeptide chain.

20 In certain embodiments, a tetramer is provided, comprising a dimer of two
 Dh3CpmFc(-)(Y349C)-GDF15(N3D):Dh3CpmFc(+)(S354C) heterodimers in which the two
 first polypeptide chains of each heterodimer are linked via an interchain disulfide bond
 between their respective GDF15 regions.

More particularly, in a specific embodiment, the tetramer comprises:

25 (a) two Dh3CpmFc(+)(S354C) domains (one each heterodimer) comprising the
 sequence of SEQ ID NO:302,

(b) two Dh3CpmFc(-)(Y349C) domains (one each heterodimer) comprising the
 sequence of SEQ ID NO:227, and

30 (c) two GDF15(N3D) polypeptides (one each heterodimer) comprising the sequence
 of SEQ ID NO:52.

In a preferred embodiment, the first polypeptide chain comprises the amino acid
 sequence:

GPSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVDGVEVHN
 AKTKPREEQYNSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPAPIEKTI
 35 SKAKGQPREPQVCTLPSSREEMTKNQVSLTCLVKGFYPSDIAVEWESNGQ
 PENNYDTTPPVLDSDGSFFLYSGLTVDKSRWQQGNVFSQVMHEALHNHY

TQKSLSLSPGARDGDHCPLGPGRCCRLHTVRASLEDLGWADWVLSPREVQ
VTMCIGACPSQFRAANMHAQIKTSLHRLKPDTPVAPCCVPASYNPMVLIQ
KTDGTGVS LQTYDDLLAKDCHCI (SEQ ID NO: 232),

which is encoded by the nucleic acid sequence:

5 ggcccgctcagtcttctcttccccccaaaaccca
 aggacaccctcatgatctcccgaccctgaggtcacatgcgtggtggtg
 gacgtgagccacgaagaccctgaggtcaagttcaactggtacgtggacgg
 cgtggaggtgcataatgccaaagacaaagccgcgggaggagcagtacaaca
 gcacgtaccgtgtggtcagcgtcctcaccgtcctgcaccaggactggctg
10 aatggcaaggagtacaagtgaaggtctccaacaaagccctcccagcccc
 catcgagaaaaccatctccaaagccaaagggcagccccgagaaccacagg
 tgtgcaccctgcccccatcccgaggagatgaccaagaaccagggtcagc
 ctgacctgcctggtcaaaggcttctatcccagcgacatcgccgtggagtg
 ggagagcaatgggcagccggagaacaactacgacaccacgcctcccgtgc
15 tggactccgacgggtccttcttctctatagcgacctcaccgtggacaag
 agcaggtggcagcaggggaacgtcttctcatgctccgtgatgcatgaggc
 tctgcacaaccactacacgcagaagagcctctccctgtctccgggtgcgc
 gcgacggagaccactgtccgctcgggccccggcggttgctgccgtctgcac
 acggtccgcgcgtcgtggaagacctgggctgggccgattgggtgctgtc
20 gccacgggaggtgcaagtgacctgtgcatcggcgcgtgcccagaccagt
 tccgggcggaacatgcacgcgcagatcaagacgagcctgcaccgcctg
 aagcccgacacggtgccagcgccctgctgcgtgcccgccagctacaatcc
 catggtgctcattcaaaagaccgacaccggggtgtcgtccagacctatg
 atgacttgtagccaaagactgccactgcata (SEQ ID NO:231).

25 In an embodiment employing the VKI signal sequence, in a preferred embodiment,
the first polypeptide chain comprises the amino acid sequence (signal sequence single
underlined):

MDMRVPAQLLGLLLLWLRGARGPSVFLFPPKPKDTLMISRTPEVTCVVV
DVSHEDPEVKFNWYVDGVEVHNAKTKPREEQYNSTYRVVSVLTVLHQDWL
30 NGKEYKCKVSNKALPAPIEKTISKAKGQPREPQVCTLPPSREEMTKNQVS
 LTCLVKGFYPSDIAVEWESNGQPENNYDTPPVLDSDGSFFLYSDLTVDK
 SRWQQGNVFSCSVMHEALHNHYTQKSLSLSPGARDGDHCPLGPGRCCRLH
 TVRASLEDLGWADWVLSPREVQVTMCIGACPSQFRAANMHAQIKTSLHRL
 KPDTVPAPCCVPASYNPMVLIQKTDGTGVS LQTYDDLLAKDCHCI (SEQ ID NO:265),

which is encoded by the nucleic acid sequence (signal sequence underlined):

atggacatgaggggtgcccgctcagctcctggggctcctgctgctgtggct
gagaggtgcgcgctgtggcccgctcagtccttcctcttccccccaaaaccca
 aggcacacctcatgatctcccgacccctgaggtcacatgctgtggtggtg
 5 gacgtgagccacgaagaccctgaggtcaagttcaactggtacgtggacgg
 cgtggaggtgcataatgccaaagacaaagccgcgggaggagcagtacaaca
 gcacgtaccgtgtggtcagcgtcctcaccgtcctgcaccaggactggctg
 aatggcaaggagtacaagtgaaggtctccaacaaagccctcccagcccc
 catcgagaaaaccatctccaaagccaaagggcagccccgagaaccacagg
 10 tgtgcaccctgcccccatccgggaggagatgaccaagaaccagggtcagc
 ctgacctgcctggtcaaaggcttctatcccagcgacatcgccgtggagtg
 ggagagcaatgggcagccggagaacaactacgacaccacgcctcccgtgc
 tggactccgacgggtccttcttctctatagcgacctcaccgtggacaag
 agcaggtggcagcaggggaacgtcttctcatgctccgtgatgcatgaggc
 15 tctgcacaaccactacacgcagaagagcctctccctgtctccgggtgcgc
 gcgacggagaccactgtccgctcgggcccgggcgttgctgccgtctgcac
 acggtccgcgcgtcgtggaagacctgggctgggccgattgggtgctgtc
 gccacgggaggtgcaagtgaccatgtgcatcggcgcgtgcccgagccagt
 tccgggcggaacatgcacgcgcagatcaagacgagcctgcaccgcctg
 20 aagcccgacacggtgccagcgccctgctgcgtgcccgccagctacaatcc
 catggtgctcattcaaaagaccgacaccggggtgtcgctccagacctatg
 atgacttggttagccaaagactgccactgcata (SEQ ID NO:266).

In a preferred embodiment, the second polypeptide chain comprises the amino acid
 sequence of SEQ ID NO:226, which is encoded by the nucleic acid sequence of SEQ ID
 25 NO:230.

In an embodiment employing the VK1 signal sequence, in a preferred embodiment,
 the second polypeptide chain comprises the amino acid sequence of SEQ ID NO:263, which
 is encoded by the nucleic acid sequence of SEQ ID NO:264.

As discussed above, a tetramer is provided comprising two polypeptide chains
 30 comprising the sequence of SEQ ID NO:232 and two polypeptide chains comprising the
 sequence of SEQ ID NO:226.

II.H.34 DhMonoFc(N297G)-GDF15

The designation "DhMonoFc(N297G)-GDF15" in the instant disclosure refers to a
 35 fusion protein comprising a GDF15 polypeptide, the N-terminus of which is linked directly to
 the C-terminus of a DhMonoFc(N297G) domain by a peptide bond.

In certain embodiments, a homodimer is provided comprising two such fusion proteins linked via interchain disulfide bond between their respective GDF15 polypeptides.

More particularly, in a specific embodiment, the homodimer comprises:

(a) two DhMonoFc(N297G) domains (one each monomer) comprising the sequence:

5 APELLGGPSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVD
 GVEVHNAKTKPREEQYGSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPA
 PIEKTISKAKGQPREPQVTTLPSSREEMTKNQVSLTCLVKGFYPSDIAVE
 WESNGQPENNYDTTPPVLDSDGSFFLYSDLTVDKSRWQQGNVVFSCSVMHE
 ALHNHYTQKSLSLSPG (SEQ ID NO:233);

10 and

(b) two GDF15 polypeptides (one each heterodimer) comprising the sequence of SEQ ID NO:12.

In a preferred embodiment, the fusion protein comprises the amino acid sequence:

APELLGGPSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVD
 GVEVHNAKTKPREEQYGSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPA
 PIEKTISKAKGQPREPQVTTLPSSREEMTKNQVSLTCLVKGFYPSDIAVE
 WESNGQPENNYDTTPPVLDSDGSFFLYSDLTVDKSRWQQGNVVFSCSVMHE
 ALHNHYTQKSLSLSPGARNGDHCPLGPGRCRLHTVRASLEDLGWADWVL
 SPREVQVTCIGACPSQFRAANMHAQIKTSLHRLKPDTPAPCCVPASYN
 PMVLIQKTDGTGVSLLQTYDDLAKDCHCI (SEQ ID NO:235),

which is encoded by the nucleic acid sequence:

gcacctgaactcctggggggaccgtcagtccttcctcttccccc
 caaaacccaaggacaccctcatgatctcccgaccctgaggtcacatgc
 gtggtggtggacgtgagccacgaagaccctgaggtcaagttcaactggtg
 25 cgtggacggcgtggaggtgcataatgccaagacaaagccgcgggaggagc
 agtacggcagcacgtaccgtgtggtcagcgtcctcaccgtcctgcaccag
 gactggctgaatggcaaggagtacaagtgaaggtctccaacaaagccct
 cccagcccccatcgagaaaaccatctccaaagccaaagggcagccccgag
 aaccacaggtgaccaccctgcccccatcccgaggagatgaccaagaac
 30 caggtcagcctgacctgcctgggtcaaaggcttctatcccagcgacatcgc
 cgtggagtgaggagcaatgggcagccggagagaacaactacgacaccacgc
 ctcccgctgctggactccgacggctccttcttctctatagcgacctcacc
 gtggacaagagcaggtggcagcaggggaacgtcttctcatgctccgtgat
 gcatgaggctctgcacaaccactacacgcagaagagcctctccctgtctc

cggggtgcgcgcaacggagaccactgtccgctcgggccccgggcgttgctgc
 cgtctgcacacgggtccgcgcgtcgtggaagacctgggctgggcccattg
 ggtgctgtcgccacgggaggtgcaagtgacctgtgcatcggcgcgtgcc
 cgagccagttccgggcggaacatgcacgcgcagatcaagacgagcctg
 5 caccgcctgaagccccgacacgggtgccagcgccctgctgctgcccgcag
 ctacaatcccatggtgctcattcaaaagaccgacaccgggggtgctcgtcc
 agacctatgatgacttgtagccaaagactgccactgcata (SEQ ID NO:234) .

As discussed above, in a specific embodiment, a homodimer is provided comprising two monomers having the sequence of SEQ ID NO:235.

10

II.H.35 DhMonoFc(N297G)-(G₄S)₄-GDF15

The designation “DhMonoFc(N297G)-(G₄S)₄-GDF15” in the instant disclosure refers to a fusion protein comprising a GDF15 polypeptide linked to a DhMonoFc(N297G) domain via a linker comprising the sequence of SEQ ID NO:18 that connects the N-terminus of the
 15 GDF15 polypeptide to the C-terminus of the DhMonoFc(N297G) domain by a peptide bond.

In certain embodiments, a homodimer is provided comprising two such fusion proteins linked via interchain disulfide bond between their respective GDF15 polypeptides.

More particularly, in a specific embodiment, the homodimer comprises:

(a) two DhMonoFc(N297G) domains (one each monomer) comprising the sequence:

20 APELLGGPSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVD
 GVEVHNAKTKPREEQYGSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPA
 PIEKTISKAKGQPREPQVTTLPSSREEMTKNQVSLTCLVKGFYPSDIAVE
 WESNGQPENNYDTPPVLDSDGSFFLYSDLTVDKSRWQQGNVVFSCSVME
 ALHNHYTQKSLSLSPG (SEQ ID NO:236) ;

25 (b) two GDF15 polypeptides (one each monomer) comprising the sequence of SEQ ID NO:12; and

(c) two polypeptide linkers (one each monomer) comprising the sequence of SEQ ID NO:18 each linking the N-terminus of a GDF15 polypeptide to the C-terminus of a DhMonoFc(N297G) domain via peptide bonds.

30 In a preferred embodiment, the fusion protein comprises the amino acid sequence (linker double underlined):

APELLGGPSVFLFPPKPKDTLMISRTPEVTCVVVDVSHEDPEVKFNWYVD
 GVEVHNAKTKPREEQYGSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPA
 PIEKTISKAKGQPREPQVTTLPSSREEMTKNQVSLTCLVKGFYPSDIAVE
 35 WESNGQPENNYDTPPVLDSDGSFFLYSDLTVDKSRWQQGNVVFSCSVME

ALHNHYTQKSLSLSPGGGGSGGGGSGGGGSGGGGSARNGDHCPLGPGRG
 CRLHTVRASLEDLGWADWVLSPREVQVTCIGACPSQFRAANMHAQIKTS
 LHRLKPDTVPAFCCVPASYNPMVLIQKTDGTGVSQTYDDLLAKDCHCI (SEQ ID
 NO: 238),

5 which is encoded by the nucleic acid sequence:

gcacctgaactcctggggggaccggtcagtccttcctcttcccc
 caaaacccaaggacaccctcatgatctcccgaccctgaggtcacatgc
 gtggtggtggacgtgagccacgaagaccctgaggtcaagttcaactggtg
 cgtggacggcgtggaggtgcataatgccaagacaaagccgaggaggagc
 10 agtacggcagcacgtaccgtgtggtcagcgtcctcacctcctgcaccag
 gactggctgaatggcaaggagtacaagtgaaggtctccaacaaagccct
 cccagcccccatcgagaaaaccatctccaaagccaaagggcagccccgag
 aaccacaggtgaccaccctgcccccatcccgaggagatgaccaagaac
 caggtcagcctgacctgcctggtcaaaggcttctatcccagcgacatcgc
 15 cgtggagtgaggagcaatgggcagccggagaacaactacgacaccacgc
 ctcccgctgctggactccgacggctccttcttctctatagcgacctcacc
 gtggacaagagcaggtggcagcaggggaacgtcttctcatgctccgtgat
 gcatgaggctctgcacaaccactacacgcagaagagcctctccctgtctc
 cgggtggaggtggtggatccggaggcgggtggaagcggaggtggtggatct
 20 ggaggcgggtggaagcgcgcgcaacggagaccactgtccgctcgggcccgg
 gcgttgctgccgtctgcacacgggtccgcgcgtcgtggaagacctgggct
 gggccgattgggtgctgtgcgcacgggaggtgcaagtgacctgtgcac
 ggcgcgtgcccagccagttccgggcggcaaacatgcacgcgcagatcaa
 gacgagcctgcaccgcctgaagcccgcacacgggtgccagcgccctgctgcg
 25 tgcccgcagctacaatcccatggtgctcattcaaaagaccgacaccggg
 gtgtcgctccagacctatgatgacttgtagccaaagactgccactgcata (SEQ ID
 NO: 237) .

As discussed above, in a specific embodiment, a homodimer is provided comprising
 two monomers having the sequence of SEQ ID NO:238.

30

II.H.36 DhMonoFc(N297G)-G₄-GDF15

The designation “DhMonoFc(N297G)-G₄-GDF15” in the instant disclosure refers to a
 fusion protein comprising a GDF15 polypeptide linked to a DhMonoFc(N297G) domain via a
 linker comprising the sequence of SEQ ID NO:58 that connects the N-terminus of the GDF15
 35 polypeptide to the C-terminus of the DhMonoFc(N297G) domain by a peptide bond.

In certain embodiments, a homodimer is provided comprising two such fusion proteins linked via interchain disulfide bond between their respective GDF15 polypeptides.

More particularly, in a specific embodiment, the homodimer comprises:

- (a) two DhMonoFc(N297G) domains (one each monomer) comprising the sequence of SEQ ID NO:236;
- (b) two GDF15 polypeptides (one each monomer) comprising the sequence of SEQ ID NO:12; and
- (c) two polypeptide linkers (one each monomer) comprising the sequence of SEQ ID NO:58 each linking the N-terminus of a GDF15 polypeptide to the C-terminus of a DhMonoFc(N297G) domain by peptide bonds.

In a preferred embodiment, the fusion protein comprises the amino acid sequence (linker double underlined):

APELLGGPSVFLFPPKPKDTLMI SRTPEVTCVVVDVSHEDPEVKFNWYVD
 GVEVHNAKTKPREEQYGSTYRVVSVLTVLHQDWLNGKEYKCKVSNKALPA
 15 PIEKTISKAKGQPREPQVTTLPSSREEMTKNQVSLTCLVKGFYPSDIAVE
 WESNGQPENNYDTPPVLDSDGSFFLYSDLTVDKSRWQQGNVFSQSVME
 ALHNHYTQKSLSLSPGGGGGARNGDHCPLGPRCCRLHTVRASLEDLGWA
 DWVLSPREVQVTMCIGACPSQFRAANMHAQIKTSLHRLKPDTVPAPCCVP
 ASYNPMVLIQKTDGTGSLQTYDDLAKDCHCI (SEQ ID NO:240),

- 20 which is encoded by the nucleic acid sequence:

gcacctgaactcctggggggaccgtcagtccttcctcttcccc
 caaaacccaaggacaccctcatgatctcccgaccctgaggtcacatgc
 gtggtggtggacgtgagccacgaagaccctgaggtcaagttcaactggtg
 cgtggacggcgtggaggtgcataatgccaagacaaagccgaggaggagc
 25 agtacggcagcacgtaccgtgtggtcagcgtcctcaccgtcctgcaccag
 gactggctgaatggcaaggagtacaagtgaaggtctccaacaaagccct
 cccagcccccatcgagaaaaccatctccaaagccaaagggcagccccgag
 aaccacaggtgaccaccctgcccccatcccgaggagatgaccaagaac
 caggtcagcctgacctgcctgggtcaaaggcttctatcccagcgacatcgc
 30 cgtggagtgaggagcaatgggcagccggagaaactacgacaccacgc
 ctcccgctgctggactccgacggctccttcttctctatagcgacctcacc
 gtggacaagagcaggtggcagcaggggaacgtcttctcatgctccgtgat
 gcatgaggctctgcacaaccactacacgcagaagagcctctccctgtctc
 cgggtggaggtggtggagcgcgcaacgggagaccactgtccgctcgggccc
 35 gggcggttgctgccgtctgcacacgggtccgcgcgtcgctggaagacctggg

ctggggccgattgggtgctgctgccacgggaggtgcaagtgaccatgtgca
 tcggcgcggtgcccagccagttccgggcggaacatgcacgcgcagatc
 aagacgagcctgcaccgcctgaagccccgacacgggtgccagcgccctgctg
 cgtgcccggcagctacaatcccatgggtgctcattcaaaagaccgacaccg
 5 ggggtgctgctccagacctatgatgaacttgtagccaaagactgccactgc
 ata (SEQ ID NO:239).

As discussed above, in a specific embodiment, a homodimer is provided comprising two monomers having the sequence of SEQ ID NO:240.

10 **III. GDF15 Polypeptides and Constructs Comprising GDF15, Including Mutant Forms Thereof**

As disclosed herein, the GDF15 polypeptides (including the full length and mature forms of human GDF15) and the constructs comprising GDF15 described in the instant disclosure can be engineered and/or produced using standard molecular biology methodology
 15 to form a mutant form of the GDF15 polypeptides and constructs provided herein. In various examples, a nucleic acid sequence encoding a mutant form of the GDF15 polypeptides and constructs provided herein, which can comprise all or a portion of SEQ ID NOs:4, 8 or 12 can be isolated and/or amplified from genomic DNA, or cDNA using appropriate oligonucleotide primers. Primers can be designed based on the nucleic and amino acid sequences provided
 20 herein according to standard (RT)-PCR amplification techniques. The amplified GDF15 mutant polypeptide nucleic acid can then be cloned into a suitable vector and characterized by DNA sequence analysis.

Oligonucleotides for use as probes in isolating or amplifying all or a portion of a mutant form of the GDF15 polypeptides and constructs provided herein can be designed and
 25 generated using standard synthetic techniques, *e.g.*, automated DNA synthesis apparatus, or can be isolated from a longer sequence of DNA.

III.A. GDF15 Polypeptide and Polynucleotide Sequences

In vivo, GDF15 is expressed as a contiguous amino acid sequence comprising a signal
 30 sequence, a pro domain and an active domain.

The 308 amino acid sequence of full length human GDF15 is:

MPGQELRTVNGSQMLLVLLVLSWLPHGGLSLAEASRASFPGPSELHSED
 SRFRELRKRYEDLLTRLRANQSWEDSNTDLVPAPAVRILTPEVRLGSGGH
 LHLRISRRAALPEGLPEASRLHRALFRLSPTASRSWDVTRPLRRQLSLARP
 35 QAPALHLRLSPPPSQSDQLLAESSSARPQLELHLRPQAARGRRRARARNG

DHCPLGPGRCCRLHTVRASLEDLGWADWVLSPREVQVTMCIGACPSQFRA
 ANMHAQIKTSLHRLKPDTPAPCCVPASYNPMVLIQKTDGTGVS LQTYDDL
 LAKDCHCI (SEQ ID NO:4)

and is encoded by the DNA sequence:

5 atgccccgggcaagaactcaggacgggtgaatggctctcagatgctcctggg
 gttgctggtgctctcgtggctgccgcattggggcgccctgtctctggccg
 aggcgagccgcgcaagtttcccgaggacctcagagttgcactccgaagac
 tccagattccgagagttgcggaaacgctacgaggacctgctaaccaggct
 gcgggccaaccagagctgggaagattcgaacaccgacctcgtcccgcccc
 10 ctgcagtccggatactcacgccagaagtgcggctgggatccggcgggccac
 ctgcacctgcgtatctctcggggcgcccttcccagaggggtccccgaggc
 ctcccgcccttaccgggctctgttcgggtgtccccgacggcgctcaaggt
 cgtgggacgtgacacgacctgcggcgctcagctcagccttgcaagacct
 caggcgcccgcgctgcacctgcgaactgtcgccgcccgcgtcgcagtcgga
 15 ccaactgctggcagaatcttcgtccgcacggccccagctggagttgcact
 tgcggccgcaagccgcccagggggcgccgcagagcgctgcgcgcaacggg
 gacctgtccgctcggggccggggcggttgcgtgccgtctgcacacggtcg
 cgcgtcgttggaagacctggggtgggcccattgggtgctgtcgccacggg
 aggtgcaagtgacctgtgcatcggcgcgtgcccagccagttccggggcg
 20 gcaaacatgcacgcgcagatcaagacgagcctgcaccgcctgaagcccga
 cacgggtgccagcgccctgctgcgtgcccgccagctacaatcccatggtgc
 tcattcaaaagaccgacaccgggggtgtcgctccagacctatgatgacttg
 ttagccaaagactgccactgcatatga (SEQ ID NO:3).

The 303 amino acid sequence of full length murine GDF15 is:

25 MAPPALQAQPPGGSQLRFLFLLLLLLLLLSWPSQGDALAMPEQRPSGPES
 QLNADDELGRFQDLSRLHANQSREDSNSEPSDPVAVRILSPEVRLGSHG
 QLLLRVNRASLSQGLPEAYRVHRALLLLTPTARPWDITRPLKRALSLRGP
 RAPALRLRLTPPPDLAMLPSGGTQLELRRLRVAAGRGRSAHAHPRDSCPL
 GPGRCCHLETQATLEDLGWSDWVLSRQLQLSMCVGECPHLYRSANTHA
 30 QIKARLHGLQPDKVPAPCCVPSSYTPVVLMHRTDSGVSLQTYDDLVARGC
 HCA (SEQ ID NO:6)

and is encoded by the DNA sequence:

atggccccgcccgcgtccaggcccagcctccaggcggtctctcaactgag
 gttcctgctgttcctgctgctgttgctgctgctgctgctcatggccatcgc
 35 agggggacgccctggcaatgcctgaacagcgacctccggccctgagtcc

caactcaacgccgacgagctacggggctcgcttcaggacctgctgagccg
 gctgcatgcccaaccagagccgagaggactcgaactcagaaccaagtctctg
 acccagctgtccggatactcagtcagaggtgagattgggggtcccacggc
 cagctgctactccgcgtcaaccgggcgtcgctgagtcaggggtctccccga
 5 agcctaccgcgtgcaccgagcgtgctcctgctgacgccgacggcccggc
 cctgggacatcactaggccctgaagcgtgcgctcagcctccgggggaccc
 cgtgctcccgattacgcctgcgcctgacgccgctccggacctggctat
 gctgccctctggcggcacgcagctggaactgcgcttacgggtagccgcg
 gcagggggcgccgaagcgcgcgtgcgcacccaagagactcgtgcccactg
 10 ggtccggggcgctgctgtcacttggagactgtgcaggcaactcttgaaga
 cttgggctggagcactgggtgctgtccccgcgccagctgcagctgagca
 tgtgcgtgggcgagtgctccccacctgtatcgctccgcgaacacgcgtgcg
 cagatcaaagcacgcctgcatggcctgcagcctgacaaggtgcctgcccc
 gtgctgtgtccccctccagctacaccccgggtggttcttatgcacaggacag
 15 acagtgggtgtgtcactgcagacttatgatgacctgggtggcccggggctgc
 cactgcgcttga (SEQ ID NO:5) .

The amino acid sequence of human GDF15 following cleavage of the 29 residue signal sequence is:

LSLAEASRASFPGPSELHSEDSRFRELKRYEDLLTRLRANQSWEDSNTD
 20 LVPAPAVRIILTPEVRLGSGGHLHLRISRALPEGLPEASRLHRLFRSLP
 TARSWDVTRPLRRQLSLARPQAPALHLRLSPPPSQSDQLLAESSARPQ
 LELHLRPQAARGRRRARARNGDHCPLGPGRCCRLHTVRASLEDLGWADWV
 LSPREVQVTMCIGACPSQFRAANMHAQIKTSLHRLKPDTPAPCCVPASY
 NPMVLIQKTDGTVSLQTYDDLAKDCHCI (SEQ ID NO:8)

25 and is encoded by the DNA sequence:

ctgtctctggccgagggcagccgcgcaagtttcccgggaccctcagagtt
 gcactccgaagactccagattccgagagttgcggaaacgctacgaggacc
 tgctaaccaggctgcgggccaaccagagctgggaagattcgaacaccgac
 ctcgctcccgccccctgcagtcgggatactcacgccagaagtgcggctggg
 30 atccggcgccacactgcacctgcgtatctctcgggccgccttcccagag
 ggctccccgaggcctcccgcccttcacgggctctgttccggctgtccccg
 acggcgtcaaggtcgtgggacgtgacacgaccgctgcggcgtcagctcag
 ccttgcaagaccccaggcgcccgctgcacctgcgactgtcgccgcgcg
 cgtcgcagtcggaccaactgctggcagaatcttcgtccgcacggccccag
 35 ctggagttgcacttgcggccgcaagccgccagggggcgccgcagagcgcg

tgcgcgcaacggggaccactgtccgctcgggcccggcggttgctgccgtc
 tgcacacgggtccgcgcgtcgtggaagacctgggctgggcccattgggtg
 ctgtcgccacgggaggtgcaagtgacctgtgcatcggcgcgtgcccag
 ccagttccgggcggaacatgcacgcgcagatcaagacgagcctgcacc
 5 gctgaagcccgcacgggtgccagcgccctgctgctgcccgcagctac
 aatcccatggtgctcattcaaaagaccgacaccggggtgtcgtccagac
 ctatgatgacttgtagccaaagactgccactgcatatga (SEQ ID NO:7)

The amino acid sequence of murine GDF 15 following cleavage of the 32 residue signal sequence is:

10 SQGDALAMPEQRPSGPESQLNADELGRFQDLLSRLHANQSREDSNSEPS
 PDPAVRILSPEVRLGSHGQLLLRVNRASLSQGLPEAYRVHRALLLTPTA
 RPWDITRPLKRALSLRGPRAPALRLRLTPPDLAMLPSGGTQLELRLRVA
 AGRGRRSAHAHPRDSCPLGPGRCHLETVQATLEDLGWSDWVLSPRQLQL
 SMCVGECPHLYRSANTHAQIKARLHGLQPDKVPAPCCVPSSYTPVVLHR
 15 TDSGVSLQTYDDLVARACHCA (SEQ ID NO:10)

and is encoded by the DNA sequence:

tcgcagggggacgccctggcaatgcctgaacagcgaccctccggccctga
 gtcccaactcaacgccgacgagctacggggtcgcttcaggacctgctga
 gccggctgcatgccaaccagagccgagaggactcgaactcagaaccaagt
 20 cctgaccagctgtccggatactcagtcagaggtgagattggggtecca
 cgccagctgctactccgcgtcaaccgggcgtcgtgagtcagggctctcc
 ccgaagcctaccgcgtgcaccgagcgtgctcctgctgacgccgacggcc
 cgccccctgggacatcactaggccccctgaagcgtgcgctcagcctccgggg
 acccctgctcccgattacgcctgcgcctgacgccgcctccggacctgg
 25 ctatgctgccctctggcggcacgcagctggaactgcgcttacgggtagcc
 gccggcagggggcgccgaagcgcgcgtgacgacccaagagactcgtgcc
 actgggtccggggcgctgctgtcacttgagactgtgcaggcaactcttg
 aagacttgggctggagcgtggtgctgtccccgcgcagctgcagctg
 agcatgtgcgtgggcgagtggtccccacctgtatcgctccgcgaacacgca
 30 tgcgcagatcaaagcacgcctgcatggcctgcagcctgacaaggtgcctg
 ccccgctgctgtgtccctccagctacacccccgtggttcttatgcacagg
 acagacagtgggtgtgtcactgcagacttatgatgacctggtggccccggg
 ctgccactgcgcttga (SEQ ID NO:9)

The biologically active form of GDF15 comprises a homodimer comprising two
 35 mature GDF15 monomers, each of which comprises SEQ ID NO:12. The monomer that

homodimerizes to form the native mature human GDF15 dimer is encoded by the nucleic acid sequence:

gcgcgcaacggggaccactgtccgctcgggcccgggcgttgctgccgtct
gcacacggtccgcgcgtcgctggaagacctgggctgggccgattgggtgc
5 tgtcgccacgggaggtgcaagtgaccatgtgcatcggcgcggtgcccgagc
cagttccgggcgggcaaacatgcacgcgcagatcaagacgagcctgcaccg
cctgaagcccgacacggtgccagcgcacctgctgcgtgcccgccagctaca
atcccatggtgctcattcaaaagaccgacaccggggtgtcgctccagacc
tatgatgacttgtagccaaagactgccactgcatatga (SEQ ID NO:11)

10 and comprises the amino acid sequence:

ARNGDHCPLGPGRCCRLHTVRASLEDLGWADWVLSPREVQVTCIGACPS
QFRAANMHAQIKTSLHRLKPDTVPAPCCVPASYNPMVLIQKTDGVSQT
YDDLAKDCHCI (SEQ ID NO:12).

Thus, the “native mature human GDF15 dimer” comprises two covalently associated
15 monomers comprising SEQ ID NO:12.

The amino acid sequence of the recombinant active form of the human GDF15, which comprises a homodimer comprising nine cysteines in each monomer to form one interchain disulfide bond and four intrachain disulfide bonds (shown with an optional N-terminal methionine residue in parentheses), is:

20 (M) ARNGDHCPLGPGRCCRLHTVRASLEDLGWADWVLSPREVQVTCIGA
CPSQFRAANMHAQIKTSLHRLKPDTVPAPCCVPASYNPMVLIQKTDGVS
LQTYDDLAKDCHCI (SEQ ID NO:189)

and is encoded by the DNA sequence (shown with an optional N-terminal methionine codon in parentheses):

25 (atg) gcgcgcaacggggaccactgtccgctcgggcccgggcgttgctgc
cgtctgcacacggtccgcgcgtcgctggaagacctgggctgggccgattg
ggtgctgtcgccacgggaggtgcaagtgaccatgtgcatcggcgcggtgcc
cgagccagttccgggcgggcaaacatgcacgcgcagatcaagacgagcctg
caccgcctgaagcccgacacggtgccagcgcacctgctgcgtgcccgccag
30 ctacaatcccatggtgctcattcaaaagaccgacaccggggtgtcgctcc
agacctatgatgacttgtagccaaagactgccactgcatataa (SEQ ID
NO:190).

The amino acid sequence of the recombinant active form of the murine

GDF15, which comprises a homodimer comprising nine cysteines in each monomer to form one interchain disulfide bond and four intrachain disulfide bonds, is:

(M) SAHAHPRDSCPLGPGRCCHLETVQATLEDLGWSDWVLSRQLQLSMC
 VGECPHLYRSANTHAQIKARLHGLQPDKVPAPCCVPSSYTPVVLHRTDS
 5 GVSLQTYDDLVARGCHCA (SEQ ID NO:14)

and is encoded by the DNA sequence:

(atg) agcgcgcgatgcgcacccaagagactcgtgccactgggtccgggg
 cgctgctgtcacctggagactgtgcaggcaactcttgaagacttgggctg
 gagcgactgggtgttgtccccgcgcagctgcagctgagcatgtgcgtgg
 10 gcgagtgtccccacctgtatcgctccgcgaacacgcatgcgcagatcaaa
 gcacgcctgcatggcctgcagcctgacaaggcgctgccccgtgctgtgt
 cccctccagctacacccgggtggttcttatgcacaggacagacagtgggtg
 tgtcactgcagacttatgatgacctggtggcccggggctgccactgcgct
 tga (SEQ ID NO:13).

15 As stated herein, the term “GDF15 polypeptide” refers to a GDF polypeptide comprising the human amino acid sequences SEQ ID NOs:4, 8 and 12. The term “GDF15 mutant polypeptide,” however, encompasses polypeptides comprising an amino acid sequence that differs from the amino acid sequence of a naturally-occurring GDF polypeptide sequence, *e.g.*, SEQ ID NOs: 4, 8 and 12, by one or more amino acids, such that the sequence is at least
 20 85% identical to SEQ ID NOs: 4, 8 and 12. GDF15 polypeptides can be generated by introducing one or more amino acid substitutions, either conservative or non-conservative and using naturally or non-naturally-occurring amino acids, at particular positions of the GDF15 polypeptide, or by deleting particular residues or stretches of residues.

A “conservative amino acid substitution” can involve a substitution of a native amino
 25 acid residue (*i.e.*, a residue found in a given position of the wild-type GDF15 polypeptide sequence) with a non-native residue (*i.e.*, a residue that is not found in a given position of the wild-type GDF15 polypeptide sequence) such that there is little or no effect on the polarity or charge of the amino acid residue at that position. Conservative amino acid substitutions also encompass non-naturally-occurring amino acid residues (as defined herein) that are typically
 30 incorporated by chemical peptide synthesis rather than by synthesis in biological systems. These include peptidomimetics, and other reversed or inverted forms of amino acid moieties.

Naturally-occurring residues can be divided into classes based on common side chain properties:

- (1) hydrophobic: norleucine, Met, Ala, Val, Leu, Ile;
- 35 (2) neutral hydrophilic: Cys, Ser, Thr;

- (3) acidic: Asp, Glu;
- (4) basic: Asn, Gln, His, Lys, Arg;
- (5) residues that influence chain orientation: Gly, Pro; and
- (6) aromatic: Trp, Tyr, Phe.

5 Additional groups of amino acids can also be formulated using the principles described in, *e.g.*, Creighton (1984) *PROTEINS: STRUCTURE AND MOLECULAR PROPERTIES* (2d Ed. 1993), W.H. Freeman and Company. In some instances it can be useful to further characterize substitutions based on two or more of such features (*e.g.*, substitution with a “small polar” residue, such as a Thr residue, can represent a highly
10 conservative substitution in an appropriate context).

Conservative substitutions can involve the exchange of a member of one of these classes for another member of the same class. Non-conservative substitutions can involve the exchange of a member of one of these classes for a member from another class.

Synthetic, rare, or modified amino acid residues having known similar
15 physiochemical properties to those of an above-described grouping can be used as a “conservative” substitute for a particular amino acid residue in a sequence. For example, a D-Arg residue may serve as a substitute for a typical L-Arg residue. It also can be the case that a particular substitution can be described in terms of two or more of the above described classes (*e.g.*, a substitution with a small and hydrophobic residue means substituting one amino acid
20 with a residue(s) that is found in both of the above-described classes or other synthetic, rare, or modified residues that are known in the art to have similar physiochemical properties to such residues meeting both definitions).

Nucleic acid sequences encoding a GDF15 mutant polypeptide provided herein, including those degenerate to SEQ ID NOs: 3, 7, 11 and 15, and those encoding polypeptide
25 variants of SEQ ID NOs:4, 8 and 12, form other aspects of the instant disclosure.

III.B. Vectors Useful for Expressing GDF15 Polypeptides and Constructs Comprising GDF15, Including Mutant Forms Thereof

In order to express the nucleic acid sequences encoding a polypeptide comprising a
30 GDF15 region, the appropriate coding sequences, *e.g.*, SEQ ID NOs:3, 7 and 11, can be cloned into a suitable vector and after introduction in a suitable host, the sequence can be expressed to produce the encoded polypeptide according to standard cloning and expression techniques, which are known in the art (*e.g.*, as described in Sambrook, J., Fritsh, E. F., and Maniatis, T. *Molecular Cloning: A Laboratory Manual* 2nd, ed., Cold Spring Harbor
35 Laboratory, Cold Spring Harbor Laboratory Press, Cold Spring Harbor, N.Y., 1989). The invention also relates to such vectors comprising a nucleic acid sequence according to the invention.

A “vector” refers to a delivery vehicle that (a) promotes the expression of a polypeptide-encoding nucleic acid sequence; (b) promotes the production of the polypeptide therefrom; (c) promotes the transfection/transformation of target cells therewith; (d) promotes the replication of the nucleic acid sequence; (e) promotes stability of the nucleic acid; (f) promotes detection of the nucleic acid and/or transformed/transfected cells; and/or (g) otherwise imparts advantageous biological and/or physiochemical function to the polypeptide-encoding nucleic acid. A vector can be any suitable vector, including chromosomal, non-chromosomal, and synthetic nucleic acid vectors (a nucleic acid sequence comprising a suitable set of expression control elements). Examples of such vectors include derivatives of SV40, bacterial plasmids, phage DNA, baculovirus, yeast plasmids, vectors derived from combinations of plasmids and phage DNA, and viral nucleic acid (RNA or DNA) vectors.

A recombinant expression vector can be designed for expression of a polypeptide comprising a GDF15 region in prokaryotic (*e.g.*, *E. coli*) or eukaryotic cells (*e.g.*, insect cells, using baculovirus expression vectors, yeast cells, or mammalian cells). Representative host cells include those hosts typically used for cloning and expression, including *Escherichia coli* strains TOP10F', TOP10, DH10B, DH5a, HB101, W3110, BL21(DE3) and BL21 (DE3)pLysS, BLUESCRIPT (Stratagene), mammalian cell lines CHO, CHO-K1, HEK293, 293-EBNA pIN vectors (Van Heeke & Schuster, *J. Biol. Chem.* 264: 5503-5509 (1989); pET vectors (Novagen, Madison Wis.). Alternatively, the recombinant expression vector can be transcribed and translated *in vitro*, for example using T7 promoter regulatory sequences and T7 polymerase and an *in vitro* translation system. Preferably, the vector contains a promoter upstream of the cloning site containing the nucleic acid sequence encoding the polypeptide. Examples of promoters, which can be switched on and off, include the lac promoter, the T7 promoter, the trc promoter, the tac promoter and the trp promoter.

Thus, provided herein are vectors comprising a nucleic acid sequence encoding a polypeptide comprising a GDF15 region that facilitate the expression of the polypeptide or construct of interest. In various embodiments, the vectors comprise an operably linked nucleotide sequence which regulates the expression of a polypeptide comprising a GDF15 region. A vector can comprise or be associated with any suitable promoter, enhancer, and other expression-facilitating elements. Examples of such elements include strong expression promoters (*e.g.*, a human CMV IE promoter/enhancer, an RSV promoter, SV40 promoter, SL3-3 promoter, MMTV promoter, or HIV LTR promoter, EF1alpha promoter, CAG promoter), effective poly (A) termination sequences, an origin of replication for plasmid product in *E. coli*, an antibiotic resistance gene as a selectable marker, and/or a convenient cloning site (*e.g.*, a polylinker). Vectors also can comprise an inducible promoter as opposed to a constitutive promoter such as CMV IE. In one aspect, a nucleic acid comprising a

sequence encoding a polypeptide comprising a GDF15 region which is operatively linked to a tissue specific promoter which promotes expression of the sequence in a metabolically-relevant tissue, such as liver or pancreatic tissue is provided.

5 III.C. Host Cells

In another aspect of the instant disclosure, host cells comprising the nucleic acids and vectors disclosed herein are provided. In various embodiments, the vector or nucleic acid is integrated into the host cell genome, which in other embodiments the vector or nucleic acid is extra-chromosomal.

10 Recombinant cells, such as yeast, bacterial (*e.g.*, *E. coli*), and mammalian cells (*e.g.*, immortalized mammalian cells) comprising such a nucleic acid, vector, or combinations of either or both thereof are provided. In various embodiments, cells comprising a non-integrated nucleic acid, such as a plasmid, cosmid, phagemid, or linear expression element, which comprises a sequence coding for expression of a polypeptide comprising a GDF15
15 region.

A vector comprising a nucleic acid sequence encoding a polypeptide comprising a GDF15 region can be introduced into a host cell by transformation or by transfection. Methods of transforming a cell with an expression vector are well known.

A nucleic acid encoding a polypeptide comprising a GDF15 region can be positioned
20 in and/or delivered to a host cell or host animal via a viral vector. Any suitable viral vector can be used in this capacity. A viral vector can comprise any number of viral polynucleotides, alone or in combination with one or more viral proteins, which facilitate delivery, replication, and/or expression of the nucleic acid of the invention in a desired host cell. The viral vector can be a polynucleotide comprising all or part of a viral genome, a viral protein/nucleic acid
25 conjugate, a virus-like particle (VLP), or an intact virus particle comprising viral nucleic acids and a nucleic acid encoding a polypeptide comprising a GDF15 region. A viral particle viral vector can comprise a wild-type viral particle or a modified viral particle. The viral vector can be a vector which requires the presence of another vector or wild-type virus for replication and/or expression (*e.g.*, a viral vector can be a helper-dependent virus), such as an adenoviral
30 vector amplicon. Typically, such viral vectors consist of a wild-type viral particle, or a viral particle modified in its protein and/or nucleic acid content to increase transgene capacity or aid in transfection and/or expression of the nucleic acid (examples of such vectors include the herpes virus/AAV amplicons). Typically, a viral vector is similar to and/or derived from a virus that normally infects humans. Suitable viral vector particles in this respect, include, for
35 example, adenoviral vector particles (including any virus of or derived from a virus of the adenoviridae), adeno-associated viral vector particles (AAV vector particles) or other parvoviruses and parvoviral vector particles, papillomaviral vector particles, flaviviral

vectors, alphaviral vectors, herpes viral vectors, pox virus vectors, retroviral vectors, including lentiviral vectors.

III.D. Isolation of a GDF15 Polypeptide, Construct Comprising a GDF15 Polypeptide or a Mutant Form Thereof

A polypeptide comprising a GDF15 region can be isolated using standard protein purification methods. A polypeptide comprising a GDF15 region can be isolated from a cell that has been engineered to express a polypeptide comprising a GDF15 region, for example a cell that does not naturally express native GDF15.

Protein purification methods that can be employed to isolate polypeptide comprising a GDF15 region, as well as associated materials and reagents, are known in the art. Exemplary methods of purifying polypeptide comprising a GDF15 region are provided in the Examples herein below. Additional purification methods that may be useful for isolating polypeptide comprising a GDF15 region can be found in references such as Bootcov MR, 1997, *Proc. Natl. Acad. Sci. USA* 94:11514-9, Fairlie WD, 2000, *Gene* 254: 67-76.

IV. Pharmaceutical Compositions Comprising a GDF15 Polypeptide, Construct Comprising a GDF15 Polypeptide or a Mutant Form Thereof

Pharmaceutical compositions comprising a monomer or multimer comprising a polypeptide comprising a GDF15 region are provided. Such polypeptide pharmaceutical compositions can comprise a therapeutically effective amount of a polypeptide comprising a monomer or multimer comprising a polypeptide comprising a GDF15 region in admixture with a pharmaceutically or physiologically acceptable formulation agent selected for suitability with the mode of administration. The term “pharmaceutically acceptable carrier” or “physiologically acceptable carrier” as used herein refers to one or more formulation agents suitable for accomplishing or enhancing the delivery of a monomer or multimer comprising a polypeptide comprising a GDF15 region into the body of a human or non-human subject. The term includes any and all solvents, dispersion media, coatings, antibacterial and antifungal agents, isotonic and absorption delaying agents, and the like that are physiologically compatible. Examples of pharmaceutically acceptable carriers include one or more of water, saline, phosphate buffered saline, dextrose, glycerol, ethanol and the like, as well as combinations thereof. In some cases it will be preferable to include isotonic agents, for example, sugars, polyalcohols such as mannitol, sorbitol, or sodium chloride in a pharmaceutical composition. Pharmaceutically acceptable substances such as wetting or minor amounts of auxiliary substances such as wetting or emulsifying agents, preservatives or buffers, which enhance the shelf life or effectiveness of the monomer or multimer comprising a polypeptide comprising a GDF15 region can also act as, or form a component of, a carrier.

Acceptable pharmaceutically acceptable carriers are preferably nontoxic to recipients at the dosages and concentrations employed.

A pharmaceutical composition can contain formulation agent(s) for modifying, maintaining, or preserving, for example, the pH, osmolarity, viscosity, clarity, color, isotonicity, odor, sterility, stability, rate of dissolution or release, adsorption, or penetration of the composition. Suitable formulation agents include, but are not limited to, amino acids (such as glycine, glutamine, asparagine, arginine, or lysine), antimicrobials, antioxidants (such as ascorbic acid, sodium sulfite, or sodium hydrogen-sulfite), buffers (such as borate, bicarbonate, Tris-HCl, citrates, phosphates, or other organic acids), bulking agents (such as mannitol or glycine), chelating agents (such as ethylenediamine tetraacetic acid (EDTA)), complexing agents (such as caffeine, polyvinylpyrrolidone, beta-cyclodextrin, or hydroxypropyl-beta-cyclodextrin), fillers, monosaccharides, disaccharides, and other carbohydrates (such as glucose, mannose, or dextrans), proteins (such as free serum albumin, gelatin, or immunoglobulins), coloring, flavoring and diluting agents, emulsifying agents, hydrophilic polymers (such as polyvinylpyrrolidone), low molecular weight polypeptides, salt-forming counterions (such as sodium), preservatives (such as benzalkonium chloride, benzoic acid, salicylic acid, thimerosal, phenethyl alcohol, methylparaben, propylparaben, chlorhexidine, sorbic acid, or hydrogen peroxide), solvents (such as glycerin, propylene glycol, or polyethylene glycol), sugar alcohols (such as mannitol or sorbitol), suspending agents, surfactants or wetting agents (such as pluronics; PEG; sorbitan esters; polysorbates such as Polysorbate 20 or Polysorbate 80; Triton; tromethamine; lecithin; cholesterol or tyloxapal), stability enhancing agents (such as sucrose or sorbitol), tonicity enhancing agents (such as alkali metal halides – preferably sodium or potassium chloride – or mannitol sorbitol), delivery vehicles, diluents, excipients and/or pharmaceutical adjuvants (see, *e.g.*, REMINGTON: THE SCIENCE AND PRACTICE OF PHARMACY, 19th edition, (1995); Berge et al., J. Pharm. Sci., 6661, 1-19 (1977). Additional relevant principles, methods, and agents are described in, *e.g.*, Lieberman et al., PHARMACEUTICAL DOSAGE FORMS-DISPERSE SYSTEMS (2nd ed., vol. 3, 1998); Ansel et al., PHARMACEUTICAL DOSAGE FORMS & DRUG DELIVERY SYSTEMS (7th ed. 2000); Martindale, THE EXTRA PHARMACOPEIA (31st edition), Remington's PHARMACEUTICAL SCIENCES (16th-20th and subsequent editions); The Pharmacological Basis Of Therapeutics, Goodman and Gilman, Eds. (9th ed.--1996); Wilson and Gisvolds' TEXTBOOK OF ORGANIC MEDICINAL AND PHARMACEUTICAL CHEMISTRY, Delgado and Remers, Eds. (10th ed., 1998). Principles of formulating pharmaceutically acceptable compositions also are described in, *e.g.*, Aulton, PHARMACEUTICS: THE SCIENCE OF DOSAGE FORM DESIGN, Churchill Livingstone (New York) (1988), EXTEMPORANEOUS ORAL LIQUID DOSAGE PREPARATIONS, CSHP (1998)).

The optimal pharmaceutical composition will be determined by a skilled artisan depending upon, for example, the intended route of administration, delivery format, and desired dosage (see, *e.g.*, Remington's PHARMACEUTICAL SCIENCES, *supra*). Such compositions can influence the physical state, stability, rate of *in vivo* release, and rate of *in vivo* clearance of a GDF15 polypeptide, construct comprising a GDF15 polypeptide or a mutant form thereof.

The primary vehicle or carrier in a pharmaceutical composition can be either aqueous or non-aqueous in nature. For example, a suitable vehicle or carrier for injection can be water, physiological saline solution, or artificial cerebrospinal fluid, possibly supplemented with other materials common in compositions for parenteral administration. Neutral buffered saline or saline mixed with free serum albumin are further exemplary vehicles. Other exemplary pharmaceutical compositions comprise Tris buffer of about pH 7.0-8.5, or acetate buffer of about pH 4.0-5.5, which can further include sorbitol or a suitable substitute. In one embodiment of the present invention, compositions comprising a GDF15 polypeptide, construct comprising a GDF15 polypeptide or a mutant form thereof can be prepared for storage by mixing the selected composition having the desired degree of purity with optional formulation agents (Remington's PHARMACEUTICAL SCIENCES, *supra*) in the form of a lyophilized cake or an aqueous solution. Furthermore, a product comprising a monomer or multimer comprising a polypeptide comprising a GDF15 region can be formulated as a lyophilizate using appropriate excipients such as sucrose.

The polypeptide pharmaceutical compositions can be selected for parenteral delivery. Alternatively, the compositions can be selected for inhalation or for delivery through the digestive tract, such as orally. The preparation of such pharmaceutically acceptable compositions is within the skill of the art.

The formulation components are present in concentrations that are acceptable to the site of administration. For example, buffers are used to maintain the composition at physiological pH or at a slightly lower pH, typically within a pH range of from about 5 to about 8.

When parenteral administration is contemplated, the therapeutic compositions for use in this invention can be in the form of a pyrogen-free, parenterally acceptable, aqueous solution comprising a desired GDF15 polypeptide, construct comprising a GDF15 polypeptide or a mutant form thereof, in a pharmaceutically acceptable vehicle. A particularly suitable vehicle for parenteral injection is sterile distilled water in which a GDF15 polypeptide, construct comprising a monomer or multimer comprising a polypeptide comprising a GDF15 region is formulated as a sterile, isotonic solution, properly preserved. Yet another preparation can involve the formulation of the desired molecule with an agent, such as injectable microspheres, bio-erodible particles, polymeric compounds (such as

polylactic acid or polyglycolic acid), beads, or liposomes, that provides for the controlled or sustained release of the product which can then be delivered via a depot injection. Hyaluronic acid can also be used, and this can have the effect of promoting sustained duration in the circulation. Other suitable means for the introduction of the desired molecule include
5 implantable drug delivery devices.

In one embodiment, a pharmaceutical composition can be formulated for inhalation. For example, a monomer or multimer comprising a polypeptide comprising a GDF15 region can be formulated as a dry powder for inhalation. Inhalation solutions comprising a monomer or multimer comprising a polypeptide comprising a GDF15 region can also be formulated
10 with a propellant for aerosol delivery. In yet another embodiment, solutions can be nebulized. Pulmonary administration is further described in International Publication No. WO 94/20069, which describes the pulmonary delivery of chemically modified proteins.

It is also contemplated that certain formulations can be administered orally. In one embodiment of the present invention, a monomer or multimer comprising a polypeptide
15 comprising a GDF15 region that is administered in this fashion can be formulated with or without those carriers customarily used in the compounding of solid dosage forms such as tablets and capsules. For example, a capsule can be designed to release the active portion of the formulation at the point in the gastrointestinal tract when bioavailability is maximized and pre-systemic degradation is minimized. Additional agents can be included to facilitate
20 absorption of a monomer or multimer comprising a polypeptide comprising a GDF15 region. Diluents, flavorings, low melting point waxes, vegetable oils, lubricants, suspending agents, tablet disintegrating agents, and binders can also be employed.

Another pharmaceutical composition can involve an effective quantity of a monomer or multimer comprising a polypeptide comprising a GDF15 region in a mixture with non-
25 toxic excipients that are suitable for the manufacture of tablets. By dissolving the tablets in sterile water, or another appropriate vehicle, solutions can be prepared in unit-dose form. Suitable excipients include, but are not limited to, inert diluents, such as calcium carbonate, sodium carbonate or bicarbonate, lactose, or calcium phosphate; or binding agents, such as starch, gelatin, or acacia; or lubricating agents such as magnesium stearate, stearic acid, or
30 talc.

Additional pharmaceutical compositions comprising a monomer or multimer comprising a polypeptide comprising a GDF15 region will be evident to those skilled in the art, including formulations comprising a monomer or multimer comprising a polypeptide comprising a GDF15 region, in sustained- or controlled-delivery formulations. Techniques for
35 formulating a variety of other sustained- or controlled-delivery means, such as liposome carriers, bio-erodible microparticles or porous beads and depot injections, are also known to those skilled in the art (see, *e.g.*, International Publication No. WO 93/15722, which describes

the controlled release of porous polymeric microparticles for the delivery of pharmaceutical compositions, and Wischke & Schwendeman, 2008, Int. J. Pharm. 364: 298-327, and Freiberg & Zhu, 2004, Int. J. Pharm. 282: 1-18, which discuss microsphere/microparticle preparation and use). As described herein, a hydrogel is an example of a sustained- or controlled-delivery formulation.

Additional examples of sustained-release preparations include semipermeable polymer matrices in the form of shaped articles, e.g. films, or microcapsules. Sustained release matrices can include polyesters, hydrogels, polylactides (U.S. Patent No. 3,773,919 and European Patent No. 0 058 481), copolymers of L-glutamic acid and gamma ethyl-L-glutamate (Sidman et al., 1983, Biopolymers 22: 547-56), poly(2-hydroxyethyl-methacrylate) (Langer et al., 1981, J. Biomed. Mater. Res. 15: 167-277 and Langer, 1982, Chem. Tech. 12: 98-105), ethylene vinyl acetate (Langer et al., supra) or poly-D(-)-3-hydroxybutyric acid (European Patent No. 0 133 988). Sustained-release compositions can also include liposomes, which can be prepared by any of several methods known in the art. See, e.g., Epstein et al., 1985, Proc. Natl. Acad. Sci. U.S.A. 82: 3688-92; and European Patent Nos. 0 036 676, 0 088 046, and 0 143 949.

A pharmaceutical composition comprising a monomer or multimer comprising a polypeptide comprising a GDF15 region which is to be used for *in vivo* administration typically should be sterile. This can be accomplished by filtration through sterile filtration membranes. Where the composition is lyophilized, sterilization using this method can be conducted either prior to, or following, lyophilization and reconstitution. The composition for parenteral administration can be stored in lyophilized form or in a solution. In addition, parenteral compositions generally are placed into a container having a sterile access port, for example, an intravenous solution bag or vial having a stopper pierceable by a hypodermic injection needle.

Once the pharmaceutical composition has been formulated, it can be stored in sterile vials as a solution, suspension, gel, emulsion, solid, or as a dehydrated or lyophilized powder. Such formulations can be stored either in a ready-to-use form or in a form (e.g., lyophilized) requiring reconstitution prior to administration.

In a specific embodiment, the present invention is directed to kits for producing a single-dose administration unit. The kits can each contain both a first container having a dried protein and a second container having an aqueous formulation. Also included within the scope of this invention are kits containing single and multi-chambered pre-filled syringes (e.g., liquid syringes and lyosyringes).

The effective amount of pharmaceutical composition comprising a monomer or multimer comprising a polypeptide comprising a GDF15 region which is to be employed therapeutically will depend, for example, upon the therapeutic context and objectives. One

skilled in the art will appreciate that the appropriate dosage levels for treatment will thus vary depending, in part, upon the molecule delivered, the indication for which a monomer or multimer comprising a polypeptide comprising a GDF15 region is being used, the route of administration, and the size (body weight, body surface, or organ size) and condition (the age and general health) of the patient. Accordingly, the clinician can titer the dosage and modify the route of administration to obtain the optimal therapeutic effect. A typical dosage can range from about 0.1 µg/kg to up to about 100 mg/kg or more, depending on the factors mentioned above. In other embodiments, the dosage can range from 0.1 µg/kg up to about 100 mg/kg; or 1 µg/kg up to about 100 mg/kg; or 5 µg/kg, 10 µg/kg, 15 µg/kg, 20 µg/kg, 25 µg/kg, 30 µg/kg, 35 µg/kg, 40 µg/kg, 45 µg/kg, 50 µg/kg, 55 µg/kg, 60 µg/kg, 65 µg/kg, 70 µg/kg, 75 µg/kg, 100 µg/kg, 200 µg/kg or up to about 10 mg/kg.

The frequency of dosing will depend upon the pharmacokinetic parameters of the a monomer or multimer comprising a polypeptide comprising a GDF15 region in the formulation being used. Typically, a clinician will administer the composition until a dosage is reached that achieves the desired effect. The composition can therefore be administered as a single dose, as two or more doses (which may or may not contain the same amount of the desired molecule) over time, or as a continuous infusion via an implantation device or catheter. Further refinement of the appropriate dosage is routinely made by those of ordinary skill in the art and is within the ambit of tasks routinely performed by them. Appropriate dosages can be ascertained through use of appropriate dose-response data.

The route of administration of the pharmaceutical composition is in accord with known methods, *e.g.*, orally; through injection by intravenous, intraperitoneal, intracerebral (intraparenchymal), intracerebroventricular, intramuscular, intraocular, intraarterial, intraportal, or intralesional routes; by sustained release systems (which may also be injected); or by implantation devices. Where desired, the compositions can be administered by bolus injection or continuously by infusion, or by implantation device.

Alternatively or additionally, the composition can be administered locally via implantation of a membrane, sponge, or other appropriate material onto which the desired molecule has been absorbed or encapsulated. Where an implantation device is used, the device can be implanted into any suitable tissue or organ, and delivery of the desired molecule can be via diffusion, timed-release bolus, or continuous administration.

In order to deliver drug, *e.g.*, a monomer or multimer comprising a polypeptide comprising a GDF15 region, at a predetermined rate such that the drug concentration can be maintained at a desired therapeutically effective level over an extended period, a variety of different approaches can be employed. In one example, a hydrogel comprising a polymer such as a gelatin (*e.g.*, bovine gelatin, human gelatin, or gelatin from another source) or a

naturally-occurring or a synthetically generated polymer can be employed. Any percentage of polymer (*e.g.*, gelatin) can be employed in a hydrogel, such as 5, 10, 15 or 20%. The selection of an appropriate concentration can depend on a variety of factors, such as the therapeutic profile desired and the pharmacokinetic profile of the therapeutic molecule.

5 Examples of polymers that can be incorporated into a hydrogel include polyethylene glycol (“PEG”), polyethylene oxide, polyethylene oxide-co-polypropylene oxide, co-polyethylene oxide block or random copolymers, polyvinyl alcohol, poly(vinyl pyrrolidinone), poly(amino acids), dextran, heparin, polysaccharides, polyethers and the like.

 Another factor that can be considered when generating a hydrogel formulation is the
10 degree of crosslinking in the hydrogel and the crosslinking agent. In one embodiment, crosslinking can be achieved via a methacrylation reaction involving methacrylic anhydride. In some situations, a high degree of cross-linking may be desirable while in other situations a lower degree of crosslinking is preferred. In some cases a higher degree of crosslinking provides a longer sustained release. A higher degree of crosslinking may provide a firmer
15 hydrogel and a longer period over which drug is delivered.

 Any ratio of polymer to crosslinking agent (*e.g.*, methacrylic anhydride) can be employed to generate a hydrogel with desired properties. For example, the ratio of polymer to crosslinker can be, *e.g.*, 8:1, 16:1, 24:1, or 32:1. For example, when the hydrogel polymer is gelatin and the crosslinker is methacrylate, ratios of 8:1, 16:1, 24:1, or 32:1 methacrylic
20 anhydride:gelatin can be employed.

V. Therapeutic Uses of a GDF15 Polypeptide, Construct Comprising a GDF15 Polypeptide or a Mutant Form Thereof

 A monomer or multimer comprising a polypeptide comprising a GDF15 region can
25 be used to treat, diagnose or ameliorate, a metabolic condition or disorder. In one embodiment, the metabolic disorder to be treated is diabetes, *e.g.*, type 2 diabetes. In another embodiment, the metabolic condition or disorder is obesity. In other embodiments the metabolic condition or disorder is dyslipidemia, elevated glucose levels, elevated insulin levels or diabetic nephropathy. For example, a metabolic condition or disorder that can be
30 treated or ameliorated using a monomer or multimer comprising a polypeptide comprising a GDF15 region includes a state in which a human subject has a fasting blood glucose level of 125 mg/dL or greater, for example 130, 135, 140, 145, 150, 155, 160, 165, 170, 175, 180, 185, 190, 195, 200 or greater than 200 mg/dL. Blood glucose levels can be determined in the fed or fasted state, or at random. The metabolic condition or disorder can also comprise a
35 condition in which a subject is at increased risk of developing a metabolic condition. For a human subject, such conditions include a fasting blood glucose level of 100 mg/dL. Conditions that can be treated using a pharmaceutical composition comprising a GDF15

mutant polypeptide can also be found in the American Diabetes Association Standards of Medical Care in Diabetes Care-2011, American Diabetes Association, Diabetes Care Vol. 34, No. Supplement 1, S11-S61, 2010.

5 In application, a metabolic disorder or condition, such as Type 2 diabetes, elevated glucose levels, elevated insulin levels, dyslipidemia, obesity or diabetic nephropathy, can be treated by administering a therapeutically effective dose of a GDF15 polypeptide, construct comprising a monomer or multimer comprising a polypeptide comprising a GDF15 region to a patient in need thereof. The administration can be performed as described herein, such as by IV injection, intraperitoneal (IP) injection, subcutaneous injection, intramuscular injection, or
10 orally in the form of a tablet or liquid formation. In some situations, a therapeutically effective or preferred dose of a monomer or multimer comprising a polypeptide comprising a GDF15 region can be determined by a clinician. A therapeutically effective dose of a monomer or multimer comprising a polypeptide comprising a GDF15 region will depend, *inter alia*, upon the administration schedule, the unit dose of agent administered, whether the
15 a monomer or multimer comprising a polypeptide comprising a GDF15 region is administered in combination with other therapeutic agents, the immune status and the health of the recipient. The term "therapeutically effective dose," as used herein, means an amount of a monomer or multimer comprising a polypeptide comprising a GDF15 region that elicits a biological or medicinal response in a tissue system, animal, or human being sought by a
20 researcher, medical doctor, or other clinician, which includes alleviation or amelioration of the symptoms of the disease or disorder being treated, *i.e.*, an amount of a monomer or multimer comprising a polypeptide comprising a GDF15 region that supports an observable level of one or more desired biological or medicinal response, for example, lowering blood glucose, insulin, triglyceride, or cholesterol levels; reducing body weight; or improving
25 glucose tolerance, energy expenditure, or insulin sensitivity.

It is noted that a therapeutically effective dose of a monomer or multimer comprising a polypeptide comprising a GDF15 region can also vary with the desired result. Thus, for example, in situations in which a lower level of blood glucose is indicated a dose of a monomer or multimer comprising a polypeptide comprising a GDF15 region will be
30 correspondingly higher than a dose in which a comparatively lower level of blood glucose is desired. Conversely, in situations in which a higher level of blood glucose is indicated at a dose of a monomer or multimer comprising a polypeptide comprising a GDF15 region will be correspondingly lower than a dose in which a comparatively higher level of blood glucose is desired.

35 In various embodiments, a subject is a human having a blood glucose level of 100 mg/dL or greater can be treated with a monomer or multimer comprising a polypeptide comprising a GDF15 region.

In one embodiment, a method of the instant disclosure comprises first measuring a baseline level of one or more metabolically-relevant compounds such as glucose, insulin, cholesterol, lipid in a subject. A pharmaceutical composition comprising a monomer or multimer comprising a polypeptide comprising a GDF15 region is then administered to the subject. After a desired period of time, the level of the one or more metabolically-relevant compounds (*e.g.*, blood glucose, insulin, cholesterol, lipid) in the subject is again measured. The two levels can then be compared in order to determine the relative change in the metabolically-relevant compound in the subject. Depending on the outcome of that comparison another dose of the pharmaceutical composition comprising a monomer or multimer comprising a polypeptide comprising a GDF15 region can be administered to achieve a desired level of one or more metabolically-relevant compound.

It is noted that a pharmaceutical composition comprising a monomer or multimer comprising a polypeptide comprising a GDF15 region can be co-administered with another compound. The identity and properties of compound co-administered with the a monomer or multimer comprising a polypeptide comprising a GDF15 region will depend on the nature of the condition to be treated or ameliorated. A non-limiting list of examples of compounds that can be administered in combination with a pharmaceutical composition comprising a monomer or multimer comprising a polypeptide comprising a GDF15 region include rosiglitazone, pioglitazone, repaglinide, nateglinide, metformin, exenatide, stiaagliptin, pramlintide, glipizide, glimepirideacarbose, and miglitol.

VI. Kits

Also provided are kits for practicing the disclosed methods. Such kits can comprise a pharmaceutical composition such as those described herein, including nucleic acids encoding the peptides or proteins provided herein, vectors and cells comprising such nucleic acids, and pharmaceutical compositions comprising such nucleic acid-containing compounds, which can be provided in a sterile container. Optionally, instructions on how to employ the provided pharmaceutical composition in the treatment of a metabolic disorder can also be included or be made available to a patient or a medical service provider.

In one aspect, a kit comprises (a) a pharmaceutical composition comprising a therapeutically effective amount of a monomer or multimer comprising a polypeptide comprising a GDF15 region; and (b) one or more containers for the pharmaceutical composition. Such a kit can also comprise instructions for the use thereof; the instructions can be tailored to the precise metabolic disorder being treated. The instructions can describe the use and nature of the materials provided in the kit. In certain embodiments, kits include instructions for a patient to carry out administration to treat a metabolic disorder, such as

elevated glucose levels, elevated insulin levels, obesity, type 2 diabetes, dyslipidemia or diabetic nephropathy.

Instructions can be printed on a substrate, such as paper or plastic, etc, and can be present in the kits as a package insert, in the labeling of the container of the kit or components thereof (*e.g.*, associated with the packaging), etc. In other embodiments, the instructions are present as an electronic storage data file present on a suitable computer readable storage medium, *e.g.* CD-ROM, diskette, etc. In yet other embodiments, the actual instructions are not present in the kit, but means for obtaining the instructions from a remote source, such as over the internet, are provided. An example of this embodiment is a kit that includes a web address where the instructions can be viewed and/or from which the instructions can be downloaded.

Often it will be desirable that some or all components of a kit are packaged in suitable packaging to maintain sterility. The components of a kit can be packaged in a kit containment element to make a single, easily handled unit, where the kit containment element, *e.g.*, box or analogous structure, may or may not be an airtight container, *e.g.*, to further preserve the sterility of some or all of the components of the kit.

EXAMPLES

The following examples, including the experiments conducted and results achieved, are provided for illustrative purposes only and are not to be construed as limiting the present invention.

EXAMPLE 1

Preparation of Fc-GDF15 Molecules

GDF15 fusions with knob/holeFc, HemiFc, charged pair (delHinge) Fc and charged pair (delHinge) cysteine clamp Fc sequences were stably expressed in serum free, suspension adapted CHO-K1 cell line. GDF15-Fc molecules were cloned into a stable expression vector containing puromycin resistance while the Fc chains were cloned into a hygromycin containing expression vector (Selexis, Inc.). The plasmids were transfected at a 1:1 ratio using lipofectamine LTX and cells were selected 2 days post transfection in a proprietary growth media containing 10ug/mL puromycin and 600ug/mLhygromycin. Media was exchanged 2 times per week during selection. When cells reached about 90% viability, they were scaled up for a fedbatch production run. Cells were seeded at 1e6/mL in a proprietary production media and fed on days 3, 6, and 8. The conditioned medium (CM) produced by the cells was harvested on day 10 and clarified. Endpoint viabilities typically were above 90%.

The Fc-GDF15 clarified, conditioned media was purified using a two-step chromatography procedure. Approximately 5 L of the CM was applied directly to a GE MabSelect SuRe column that had previously been equilibrated with Dulbecco's Phosphate

Buffered Saline (PBS). The bound protein underwent three wash steps: first, 3 column volumes (CV) of PBS; next, 1 CV of 20 mM Tris, 100 mM sodium chloride, pH 7.4; and finally, 3 CV of 500 mM L-arginine, pH 7.5. These wash steps remove unbound or lightly bound media components and host cell impurities. The column was then re-equilibrated with 5 CV of 20 mM Tris, 100 mM sodium chloride at pH 7.4 which brings the UV absorbance back to baseline. The desired protein was eluted with 100 mM acetic acid at pH 3.6 and collected in bulk. The protein pool was quickly titrated to within a pH range of 5.0 to 5.5 with 1 M Tris-HCl, pH 9.2.

The pH adjusted protein pool was next loaded onto a GE SP Sepharose HP column that had been previously equilibrated with 20 mM MES at pH 6.0. The bound protein was then washed with 5 CV of equilibration buffer, and finally eluted over a 20 CV, 0 to 50% linear gradient from 0 to 400 mM sodium chloride in 20 mM MES at pH 6.0. Fractions were collected during the elution and analyzed by analytical size-exclusion chromatography (Superdex 200) to determine the appropriate fractions to pool for a homogeneous product. The SP HP chromatography removes product-related impurities such as free Fc, clipped species, and Fc-GDF15 multimers.

The SP HP pool was then buffer exchanged into 10 mM sodium acetate, 5% proline, pH 5.2 by dialysis. It was concentrated to approximately 15 mg/ml using the Sartorius Vivaspin 20 Ten kilo-Dalton molecular weight cut-off centrifugal device. Finally, it was sterile filtered and the resulting solution containing the purified Fc-GDF15 molecules is stored at 5° C. Final products were assessed for identity and purity using mass spectral analysis, sodium dodecyl sulfate polyacrylamide electrophoresis and size exclusion high performance liquid chromatography.

The purification method described above was employed to purify DhMonoFc-GDF15 fusion proteins. However, it was found that the addition of the H6D mutation to the DhMonoFc-GDF15 caused soluble aggregates to form in the SP elution. Therefore, the purification of the DhMonoFc-GFF15(H6D) included an additional SEC step (Superdex 200 with 20 mM phosphate, 250 mM NaCl, pH 6.8), followed by loading on Q-sepharose HP and eluting with a gradient from 0 to 0.6M NaCl in 20 mM tris, pH 8.5.

30

EXAMPLE 2

Preparation of GDF15-HSA and DhMonoFc Molecules

GDF15 fusions with HSA and DhMonoFc sequences were stably expressed in CHO-S cells (Invitrogen). For each of the constructs producing homodimers, the coding sequence was cloned into stable expression vector containing puromycin resistance (Selexis, Inc.). In the case of the HSA-(G₄S)₄-GDF15:GDF15 heterodimer, the HSA-(G₄S)₄-GDF15 fusion sequence was cloned into an expression vector containing puromycin resistance and the

GDF15 sequence was cloned into an expression vector containing hygromycin resistance. CHO-S parental cells were maintained in CD-CHO medium (Invitrogen) supplemented with 8 mM L-glutamine and were transfected with 4 µg of plasmid DNA using a Lipofectamine LTX transfection kit (Invitrogen) according to the manufacturer's instructions. In the case of the HSA-(G₄S)₄-GDF15:GDF15 heterodimer, the two plasmids were mixed at a 1:1 ratio prior to transfection. Stable cell lines were selected using 10 µg/mL of puromycin (homodimers) or 10 µg/mL of puromycin plus 400 µg/mL hygromycin (heterodimer). Upon recovery, which was defined as >90% viability using a Vi-Cell counter (Beckman Coulter), the stable CHO-S cell lines were expanded and used to seed either batch productions in shake flasks or fedbatch productions in WAVE bioreactors (GE Healthcare). Both processes were seeded at 1e6 viable cells/mL in production medium. Batch productions were harvested by centrifugation on day 6, while fedbatch productions were fed on days 3, 6, and 8. The CM produced by the cells was harvested by centrifugation on day 10 and clarified.

The HSA-GDF15 fusion proteins were purified from clarified conditioned media using two chromatographic steps. The clarified conditioned media containing the HSA-GDF15 fusion protein was applied to a Cibracon Blue Sepharose HP column that is equilibrated with 20mM Phosphate, 150mM NaCl pH 7.4. The column was next washed with equilibration buffer until a baseline ultraviolet (UV) level is obtained. Product and contaminants are eluted by a 20 mM Phosphate, 2M NaCl buffer and the elutions were collected and subsequently assayed by Coomassie-stained SDS-PAGE (sodium dodecyl sulfate polyacrylamide gel electrophoresis) to identify which eluate fractions contained a polypeptide that migrates at the predicted molecular weight of HSA-GDF15 fusion protein. Following the Blue Sepharose step, the pooled fractions containing product were dialyzed versus 10 mM tris, pH 8.0. The dialysis step allowed for the binding of HSA-GDF15 fusion protein when applied to anion exchange chromatographic resin. The final chromatography step was Q-Sepharose HP which applies a linear gradient (0 to 0.6M NaCl in 10 mM tris pH 8.0) to elute the bound fusion protein. The elution from the Q-Sepharose HP was collected as fractions and then assayed by SDS-PAGE and analytical size exclusion chromatography to determine the appropriate fractions to pool. LCMS and SDS-PAGE were run to confirm the identity of each protein. The resulting pool was buffer exchanged by dialysis into 10mM Sodium Acetate, 9% sucrose pH 4.5, sterile filtered, and finally stored at 5 C or frozen.

DhMonoFc-GDF15 fusion proteins were purified as set forth above in Example 2 for other Fc-GDF15 fusion proteins.

35

EXAMPLE 3

Suppression of Food Intake in Hyperphagic ob/ob Mice by Fc Fusion GDF15 Polypeptides and HSA Fusion GDF15 Polypeptides

GDF15 reduces food intake in hyperphagic ob/ob mice, and a food intake assay was used to evaluate efficacy of different forms of GDF15 analogs. As the half-life of human GDF15 polypeptide in mouse was observed to be approximately 3 hours, an Fc fusion strategy was used to extend protein half-life. Various multimers comprising a polypeptide comprising a GDF15 region were generated and analyzed for *in vivo* activity, by introducing the multimer into hyperphagic leptin-deficient ob/ob mice, and measuring the ability of a particular multimer comprising a polypeptide comprising a GDF15 region to suppress food intake in these animals. The multimer comprising a polypeptide comprising a GDF15 region to be tested was injected subcutaneously into a 7-8 week old ob/ob mouse (Jackson Laboratory) between 4-5pm on day 0. Animals were transferred after injection to cages where food had been premeasured, and food intake was measured between 9-10 AM the next day.

The results of representative experiments are provided in Figures 6-53. These experiments demonstrate that the described multimers comprising GDF15 regions exhibit a decrease in food intake in ob/ob mice, with greater potency than those of native mature hGDF15 homodimer.

EXAMPLE 4

Chronic Efficacy of GDF15 Constructs in DIO Mice

Certain multimers comprising GDF15 regions are administered chronically and subcutaneously into DIO mice, once per week. The constructs demonstrate efficacy in improving various metabolic parameters, including body weight, blood glucose levels and glucose tolerance, serum insulin levels, serum cholesterol levels, serum triglyceride levels and oral lipid tolerance.

EXAMPLE 5

In Vivo Activity of GDF15 Constructs

Male C57Bl/6 were fed a 60% high fat diet for 15 weeks and divided into different treatment groups for each group to have the same pretreatment body weight, glucose, insulin, triglyceride and cholesterol levels. Animals were subcutaneously dosed with proteins or vehicle buffer weekly for 5 weeks. Three different dose levels were selected for the proteins: 10, 1, 0.1nmol/kg, which are equivalent to 1.25, 0.125, 0.0125 mg/kg. Studies were carried for 5 weeks, with the last dose on day 28.

Body weight was measured weekly during the 5 weeks of treatment and drug washout. One oral glucose tolerance tests (OGTT) was performed 2 weeks after the first protein injection in animals fasted for 4 hours. Another oral glucose tolerance tests (OGTT) was performed 5 weeks after the first protein injection in animals fasted for 16 hours. In OGTT, animals were orally administered with 2g/kg glucose solution, and glucose levels

were measured by AlphaTRAK glucometer (Abbott) at 0, 15, 30, 60, 120min. Area under the curve (AUC) of the glucose levels during the OGTT were calculated to compare glucose tolerance of different treatment groups. Serum samples were collected at 3 weeks after first protein injection and used to measure insulin, triglyceride and cholesterol levels, as well as the levels of test articles. Insulin levels were measured using immunoassay kit (Alpco). Triglyceride and cholesterol levels were measured using enzymatic assays (Wako).

The results are shown in Figures 54-59 (asterisks indicate statistical significance). These experiments demonstrate that the described multimers comprising GDF15 regions reduce AUC of the glucose levels during the OGTT (Figures 55 and 59), reduce body weight (Figure 54) reduce insulin levels (Figure 56), reduce cholesterol (Figure 58) and reduce triglycerides (Figure 57).

EXAMPLE 6

Thermal Stability of GDF15 Constructs

The thermal stability of the selected GDF15 constructs was assessed by differential scanning calorimetry on a MicroCal Capillary VP-DSC system in which temperature differences between the reference and sample cell are continuously measured, and calibrated to power units. This data channel is referred to as the DP signal, or the differential power between the reference and sample cell. The unfolding of a protein molecule appears as an endothermic transition on the DSC thermogram and can be characterized by the thermal transition midpoints (T_m). The samples were heated from 10°C to 100°C at a heating rate of 60°C/hour. The pre-scan time was 15 minutes and the filtering period was 10 seconds. The concentrations used in the DSC experiments were around 1.0 mg/mL. The data analysis for baseline correction and determination of T_m values was done using MicroCal Origin 7 software.

In particular, a dimer of DhCpmFc(-)-GDF15(N3D):DhCpmFc(+) was compared to a dimer of Dh3CpmFc(-)-GDF15(N3D):DhCpmFc(+); a dimer of DhCpmFc(-)-GDF15(Ndel3):DhCpmFc(+) was compared to a dimer of Dh3CpmFc(-)-GDF15(Ndel3):DhCpmFc(+); a dimer of DhCpmFc(-)(Y349C)-GDF15(N3D):DhCpmFc(+)(S354C) was compared to a dimer of Dh3CpmFc(-)(Y349C)-GDF15(N3D):DhCpmFc(+)(S354C); and a dimer of DhCpmFc(-)(Y349C)-GDF15(Ndel3):DhCpmFc(+)(S354C) was compared to a dimer of Dh3CpmFc(-)(Y349C)-GDF15(Ndel3):DhCpmFc(+)(S354C). The results are shown in Figure 61. These experiments demonstrate that the “Dh3CpmFc” domains confer greater stability than the corresponding “DhCpmFc” domains.

EXAMPLE 7

Fcγ Receptor Binding Analysis

Selected GDF15 constructs were analyzed for their binding activity to Fcγ receptors on BIA3000. Each Fcγ receptor was captured on anti-his antibody coated CM5 surface (captured RL ~ 200 RU). The GDF15 constructs were diluted to 250 nM in sample buffer (0.1 mg/ml BSA, 0.005% P20, PBS). Each GDF15 construct was injected over anti-his antibody captured Fcγ receptor surfaces at 50 μL/min for 3 minutes. After a 5-minute dissociation in instrument running buffer (0.005% P20 in PBS), each Fcγ receptor surface was regenerated by an injection of 8 mM Glycine, pH1.5, 1M NaCl for 30 seconds, followed by an injection of 10 mM Glycine, pH1.5 for 30 seconds. The resulting sensorgrams were analyzed using BIAcore BIAEvaluation (v. 4.1). The binding response in the unit of RU was read at 10 seconds before end of injection.

In particular, FcγRI, FcγRIIIA and FcγRIIA were determined with respect to a dimer of DhCpmFc(-)-GDF15(Ndel3):DhCpmFc(+), a dimer of DhCpmFc(-)(Y349C)-GDF15(Ndel3): DhCpmFc(+)(S354C); a dimer of Dh3CpmFc(-)-GDF15(Ndel3); a dimer of Dh3CpmFc(-)(Y349C)-GDF15(Ndel3)-Dh3CpmFc(+)(S354C); a dimer of Dh3CpmFc(-)-GDF15(N3D); and a dimer of Dh3CpmFc(-)(Y349C)-GDF15(N3D):Dh3CpmFc(+)(S354C). The results are shown in Figure 60. These experiments demonstrate that the “Dh3CpmFc” domains essentially eliminate FcγRI, FcγRIIIA and FcγRIIA binding.

While the present invention has been described in terms of various embodiments, it is understood that variations and modifications will occur to those skilled in the art. Therefore, it is intended that the appended claims cover all such equivalent variations that come within the scope of the invention as claimed. In addition, the section headings used herein are for organizational purposes only and are not to be construed as limiting the subject matter described.

All references cited in this application are expressly incorporated by reference herein for any purpose.

CLAIMS

What is claimed is:

1. A fusion protein comprising a GDF15 region and an Fc domain.
2. The fusion protein of claim 1, wherein the Fc domain comprises a sequence
5 selected from the group consisting of SEQ ID NOs:16, 22, 28, 29, 33, 35, 38, 48, 85, 91, 106, 132, 141, 148, 155, 162, 169, 176, 183, 192, 199, 206, 213, 220, 227, 233, 236, 268, 275, 281, 282, 283, 284, 285, 286, 287, 288, 289, 290, 291, 292, 293, 294, 295, 296, 297, 298, 299, 300, 301 and 302.
3. The fusion protein of claim 2, wherein the GDF15 region comprises a
10 sequence selected from the group consisting of SEQ ID NOs:4, 8, 12, 25, 52 and 55.
4. The fusion protein of claim 2, wherein the GDF15 region and the Fc domain are joined by a polypeptide linker.
5. The fusion protein of claim 4, wherein the polypeptide linker is selected from
the group consisting of SEQ ID NOs:18, 30, 34, 40, 58, 61, 64, 69, 72, 75, 78, 113, 116, 119,
15 122, 125, 128.
6. The fusion protein of claim 1, wherein the fusion protein comprises a
sequence selected from the group consisting of SEQ ID NOs: 46, 24, 27, 32, 37, 20, 42, 50,
54, 57, 60, 63, 66, 68, 71, 74, 77, 82, 84, 88, 93, 96, 98, 100, 102, 104, 108, 134, 137, 139,
143, 146, 150, 153, 269, 272, 276, 279, 157, 160, 164, 167, 171, 174, 178, 181, 185, 188, 194,
20 197, 201, 204, 208, 211, 215, 218, 222, 225, 229, 232, 233, 238 and 240.
7. The fusion protein of claim 1, wherein the fusion protein comprises two or more Fc domains.
8. A dimer comprising (i) a first polypeptide chain comprising a GDF15 region and a first Fc domain, and (ii) a second polypeptide chain comprising a second Fc domain.
- 25 9. The dimer of claim 8, wherein first Fc domain comprises a sequence selected from the group consisting of SEQ ID NOs:16, 22, 28, 29, 33, 35, 38, 48, 85, 91, 106, 132, 141, 148, 155, 162, 169, 176, 183, 192, 199, 206, 213, 220, 227, 233, 236, 268, 275, 281, 282, 283, 284, 285, 286, 287, 288, 289, 290, 291, 292, 293, 294, 295, 296, 297, 298, 299, 300, 301 and 302.
- 30 10. The dimer of claim 9, wherein second Fc domain comprises a sequence selected from the group consisting of SEQ ID NOs:16, 22, 28, 29, 33, 35, 38, 48, 85, 91, 106,

132, 141, 148, 155, 162, 169, 176, 183, 192, 199, 206, 213, 220, 227, 233, 236, 268, 275, 281, 282, 283, 284, 285, 286, 287, 288, 289, 290, 291, 292, 293, 294, 295, 296, 297, 298, 299, 300, 301 and 302.

11. The dimer of claim 9, wherein the GDF15 region comprises a sequence
5 selected from the group consisting of SEQ ID NOs:4, 8, 12, 25, 52 and 55.

12. The dimer of claim 9, wherein the GDF15 region and the Fc domain are joined by a polypeptide linker.

13. The dimer of 12, wherein the polypeptide linker is selected from the group consisting of SEQ ID NOs:18, 30, 34, 40, 58, 61, 64, 69, 72, 75, 78, 113, 116, 119, 122, 125,
10 128.

14. The dimer of claim 8, wherein the fusion protein comprises a sequence selected from the group consisting of SEQ ID NOs: 46, 24, 27, 32, 37, 20, 42, 50, 54, 57, 60, 63, 66, 68, 71, 74, 77, 82, 84, 88, 93, 96, 98, 100, 102, 104, 108, 134, 137, 139, 143, 146, 150, 153, 269, 272, 276, 279, 157, 160, 164, 167, 171, 174, 178, 181, 185, 188, 194, 197, 201, 204,
15 208, 211, 215, 218, 222, 225, 229, 232, 233, 238 and 240.

15. The dimer of claim 8, wherein the first and second polypeptide chains are non-covalently associated.

16. The dimer of claim 8, wherein the first and second polypeptide chains are covalently associated.

17. The dimer of claim 16, wherein the first and second polypeptide chains are covalently associated via disulfide bonds between their respective Fc domains.
20

18. A tetramer comprising (i) a first dimer and (ii) a second dimer, wherein the first and second dimer independently comprise a dimer of claim 1 and wherein the first polypeptide chain of the first dimer is linked to the first polypeptide chain of the second dimer via an interchain disulfide bond between their respective GDF15 regions.
25

19. The tetramer of claim 18, wherein the Fc domain of the first polypeptide of the first dimer comprises a sequence selected from the group consisting of SEQ ID NOs:16, 22, 28, 29, 33, 35, 38, 48, 85, 91, 106, 132, 141, 148, 155, 162, 169, 176, 183, 192, 199, 206, 213, 220, 227, 233, 236, 268, 275, 281, 282, 283, 285, 286, 287, 288, 289, 290, 291, 292,
30 293, 294, 295, 296, 297, 298, 299, 300, 301 and 302.

20. The tetramer of claim 19, wherein the GDF15 region of the first polypeptide chain of the first dimer comprises a sequence selected from the group consisting of SEQ ID NOs:4, 8, 12, 25, 52 and 55.

21. The tetramer of claim 20, wherein the GDF15 region and the Fc domain are joined by a polypeptide linker.

22. The tetramer of claim 21, wherein the polypeptide linker is selected from the group consisting of SEQ ID NOs:18, 30, 34, 40, 58, 61, 64, 69, 72, 75, 78, 113, 116, 119, 122, 125, 128.

23. The tetramer of claim 18, wherein the Fc domain of the first polypeptide of the first dimer comprises a sequence selected from the group consisting of SEQ ID NOs: 46, 24, 27, 32, 37, 20, 42, 50, 54, 57, 60, 63, 66, 68, 71, 74, 77, 82, 84, 88, 93, 96, 98, 100, 102, 104, 108, 134, 137, 139, 143, 146, 150, 153, 269, 272, 276, 279, 157, 160, 164, 167, 171, 174, 178, 181, 185, 188, 194, 197, 201, 204, 208, 211, 215, 218, 222, 225, 229, 232, 233, 238 and 240.

24. The tetramer of claim 19, wherein the Fc domain of the first polypeptide of the second dimer comprises a sequence selected from the group consisting of SEQ ID NOs:16, 22, 28, 29, 33, 35, 38, 48, 85, 91, 106, 132, 141, 148, 155, 162, 169, 176, 183, 192, 199, 206, 213, 220, 227, 233, 236, 268, 275, 281, 282, 283, 284, 285, 286, 287, 288, 289, 290, 291, 292, 293, 294, 295, 296, 297, 298, 299, 300, 301 and 302.

25. The tetramer of claim 24,, wherein the Fc domain of the first polypeptide of the second dimer comprises a sequence selected from the group consisting of SEQ ID NOs: 46, 24, 27, 32, 37, 20, 42, 50, 54, 57, 60, 63, 66, 68, 71, 74, 77, 82, 84, 88, 93, 96, 98, 100, 102, 104, 108, 134, 137, 139, 143, 146, 150, 153, 269, 272, 276, 279, 157, 160, 164, 167, 171, 174, 178, 181, 185, 188, 194, 197, 201, 204, 208, 211, 215, 218, 222, 225, 229, 232, 233, 238 and 240.

Figure 1

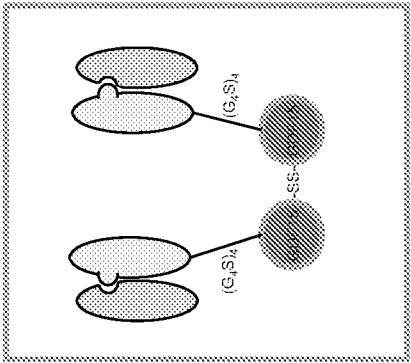


Figure 2

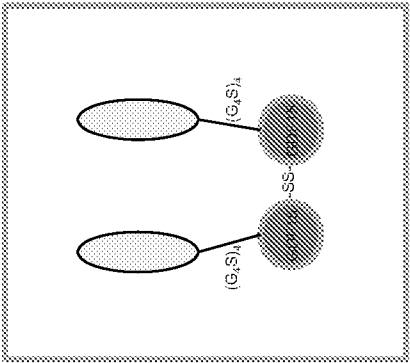


Figure 3

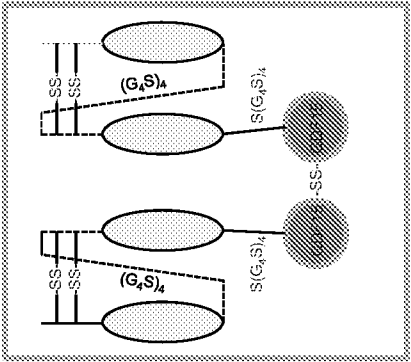


Figure 4

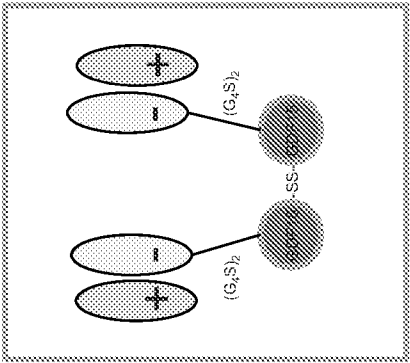


Figure 5

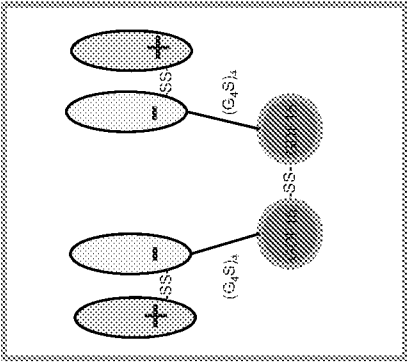


Figure 6

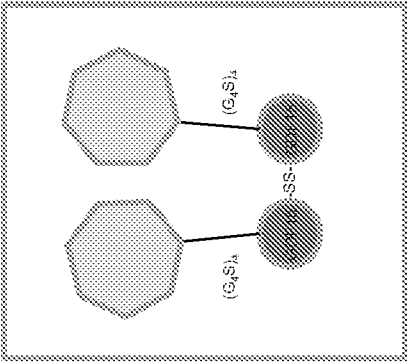


Figure 7

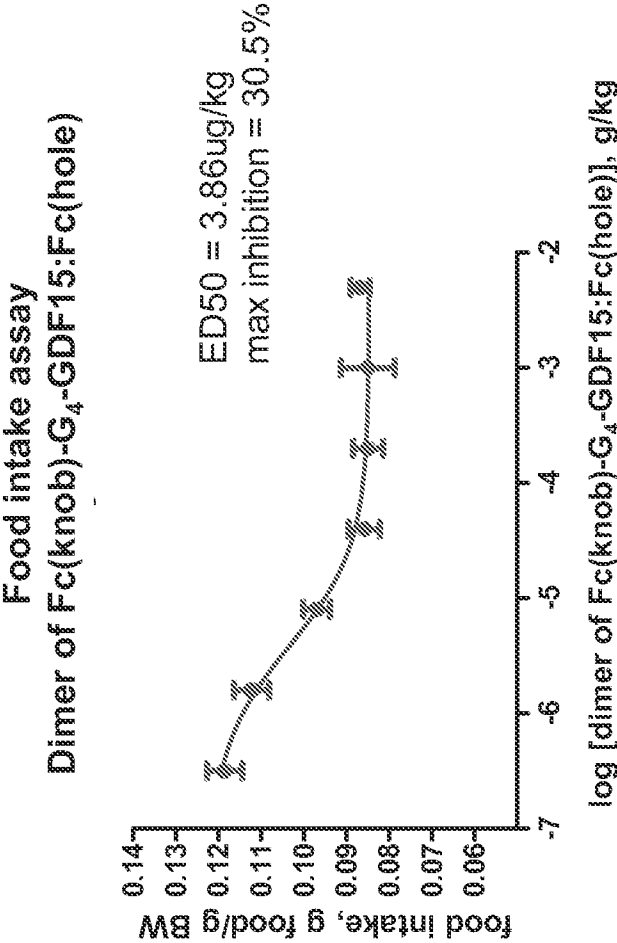
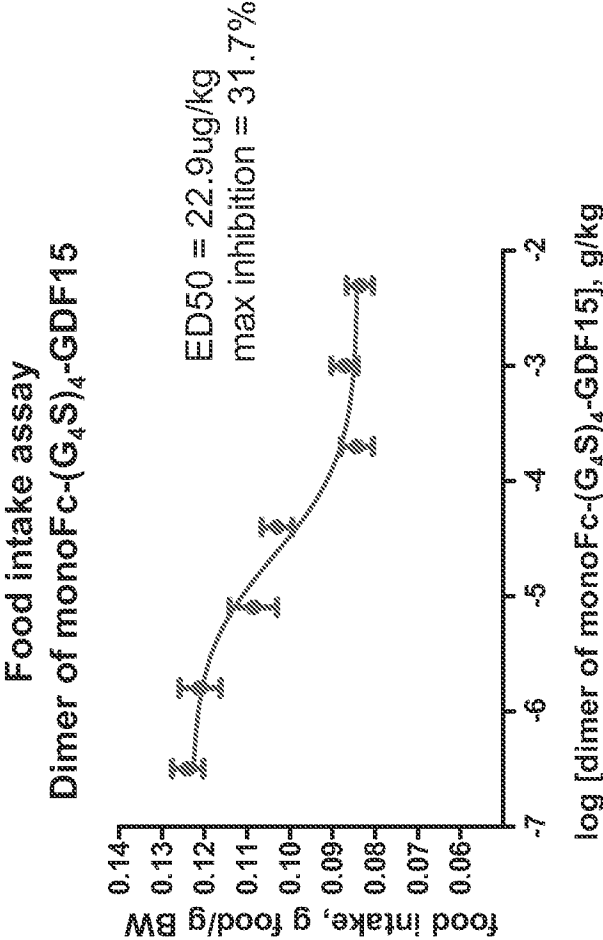
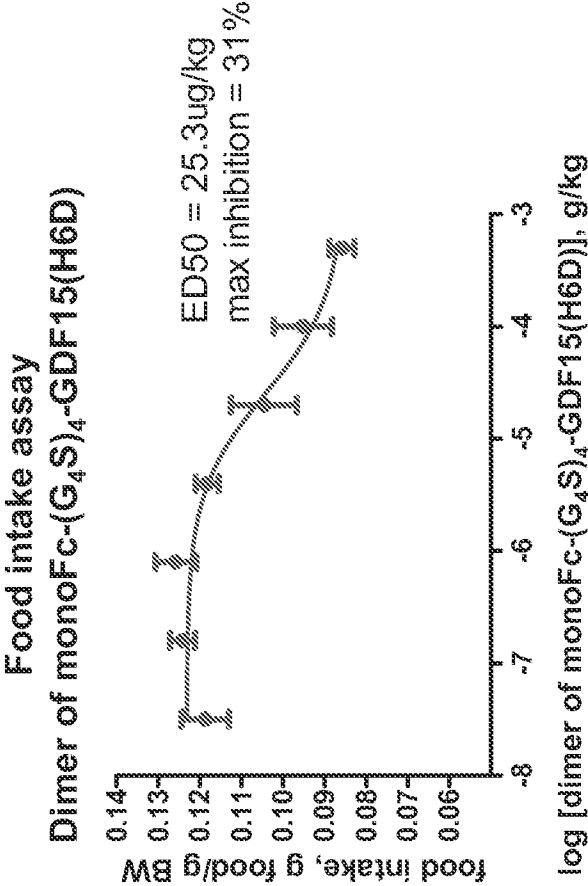


Figure 8



9/61

Figure 9



10/61

Figure 10

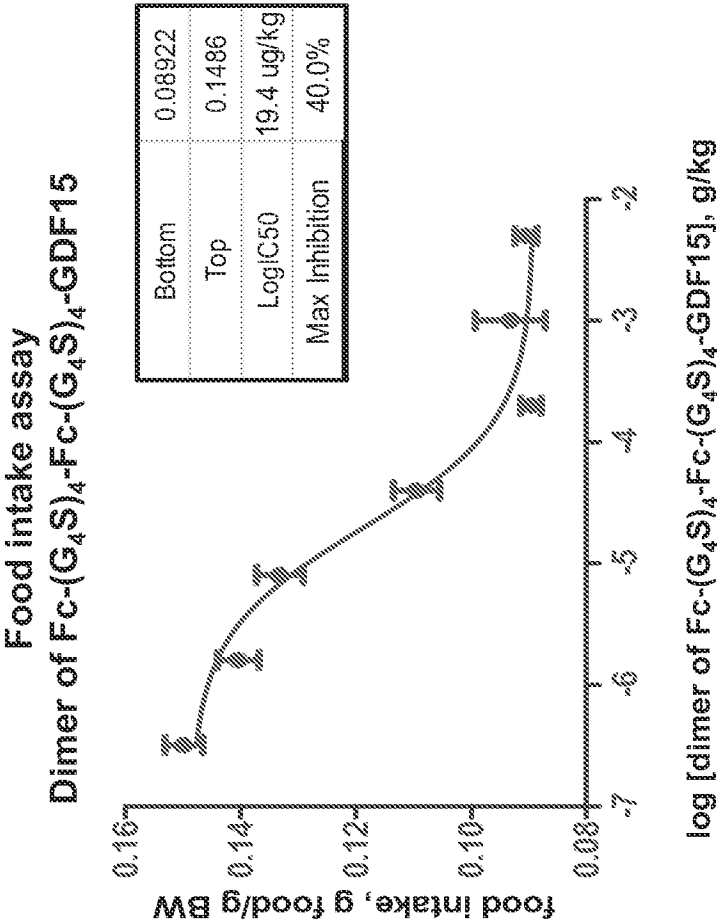
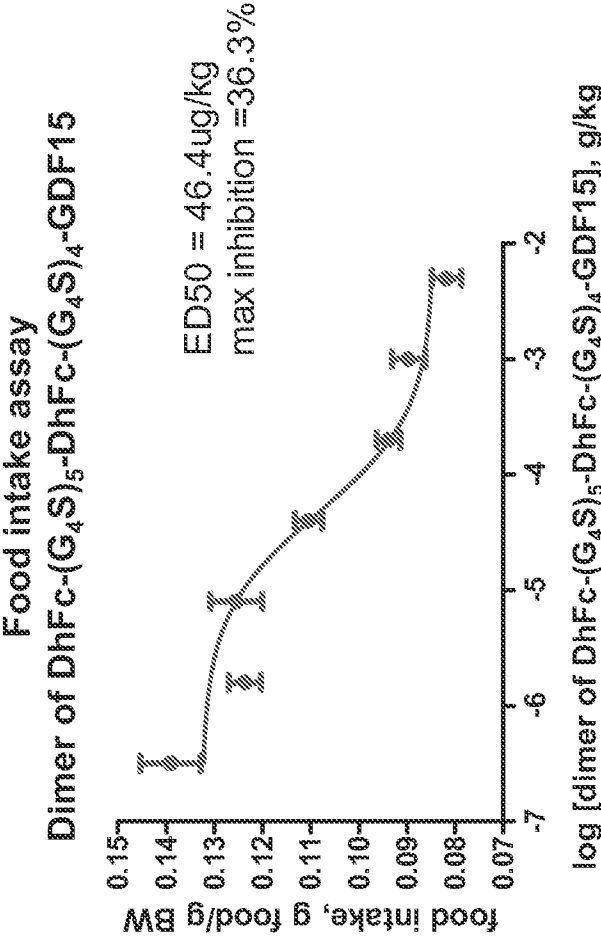


Figure 11



12/61

Figure 12

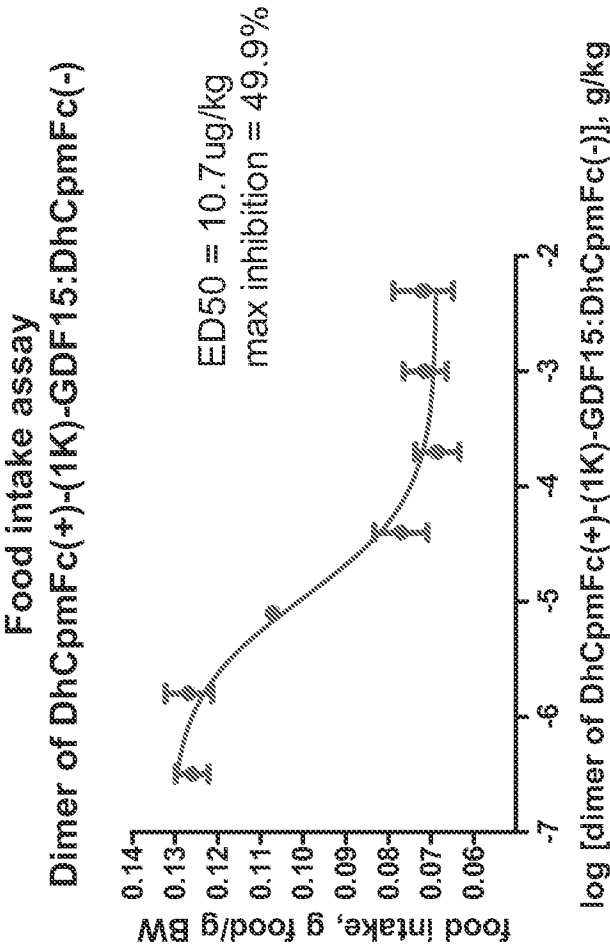


Figure 13

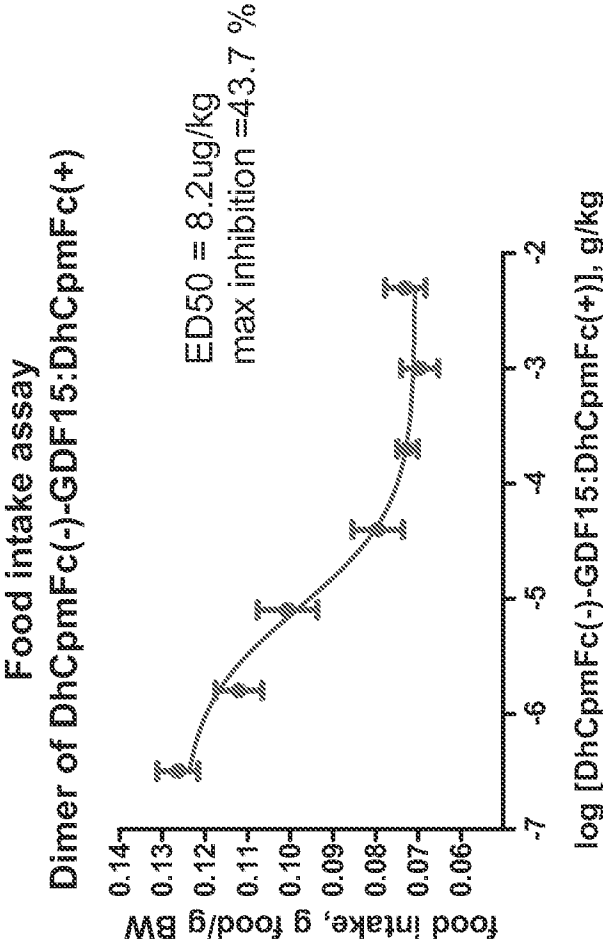


Figure 14

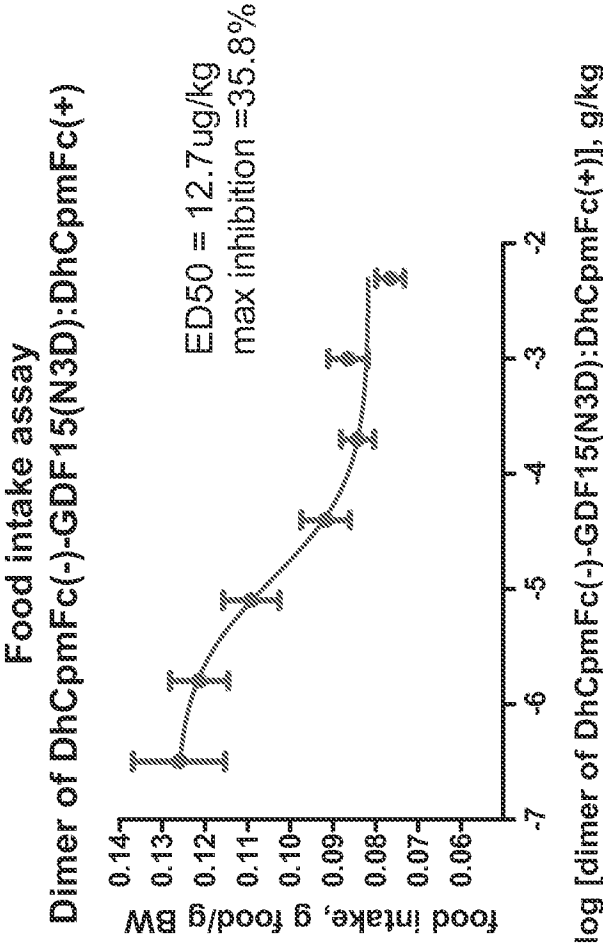


Figure 15

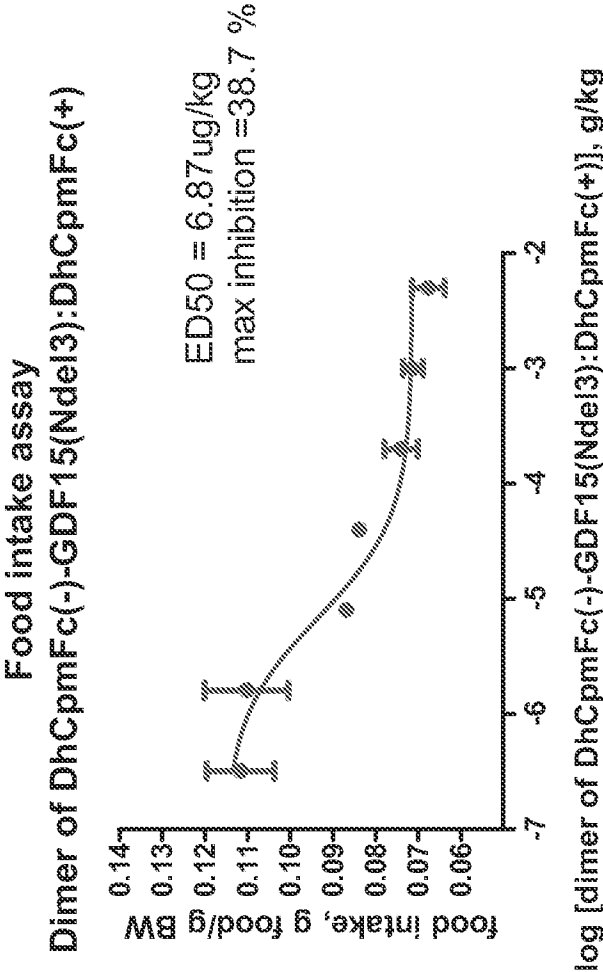
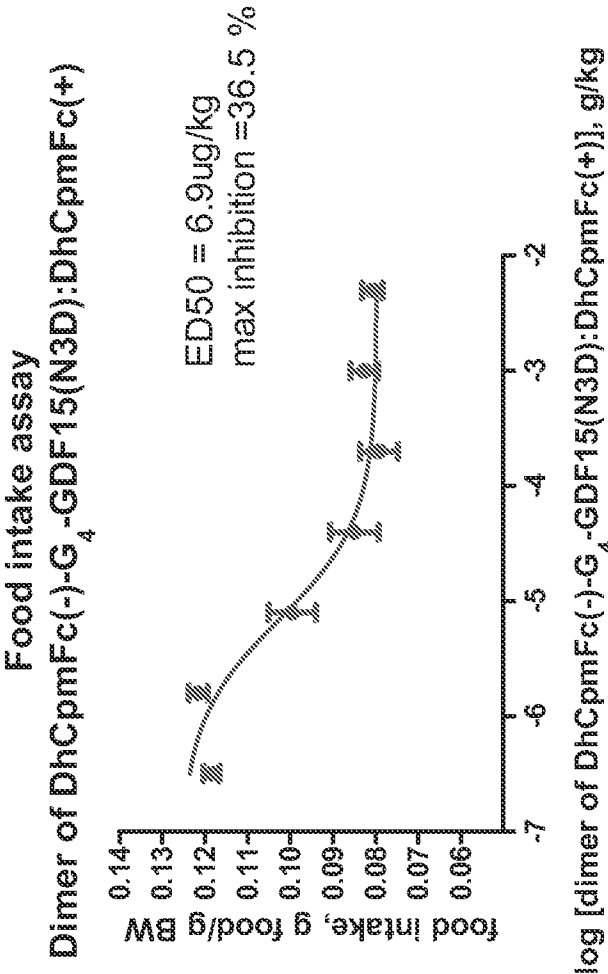
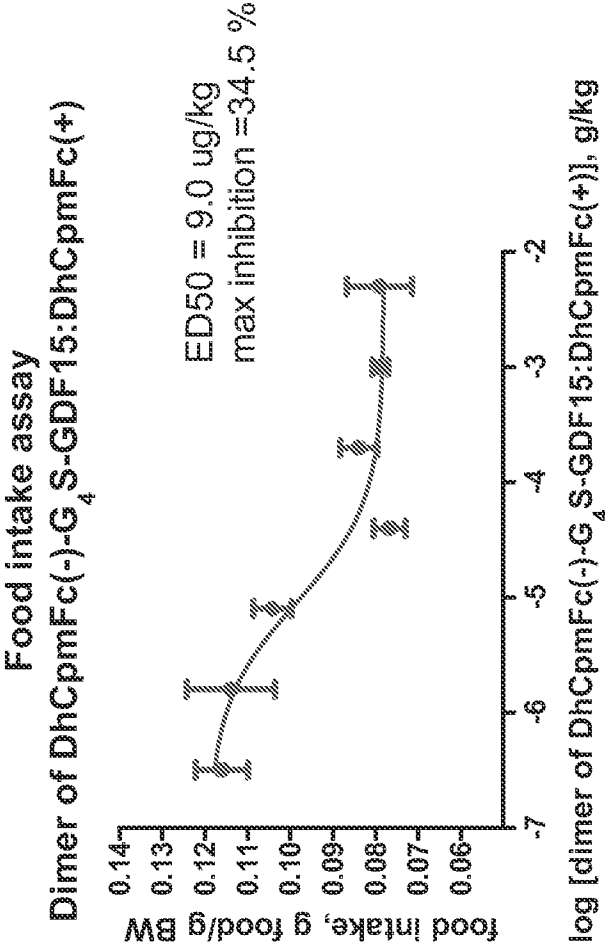


Figure 16



17/61

Figure 17



18/61

Figure 18

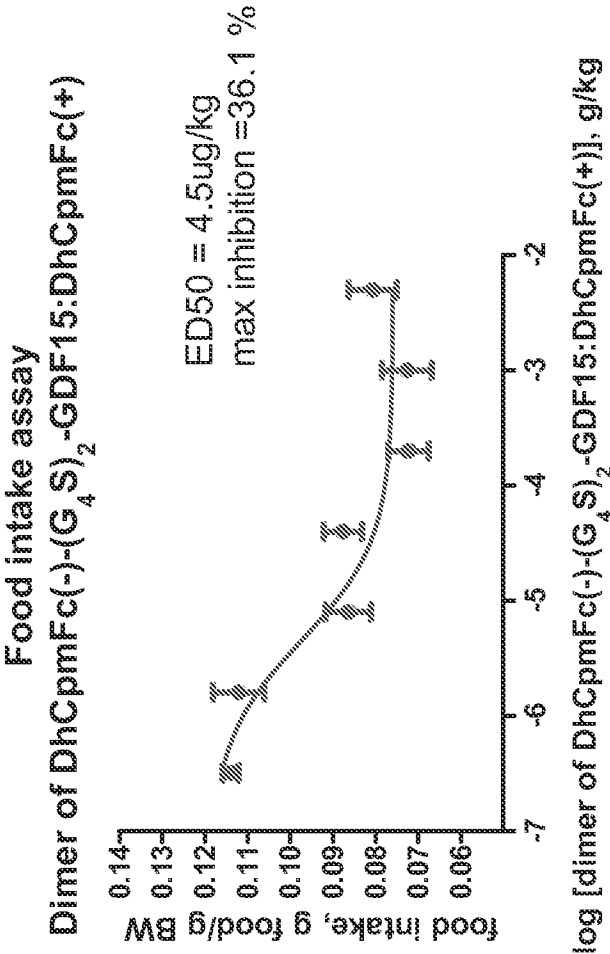


Figure 19

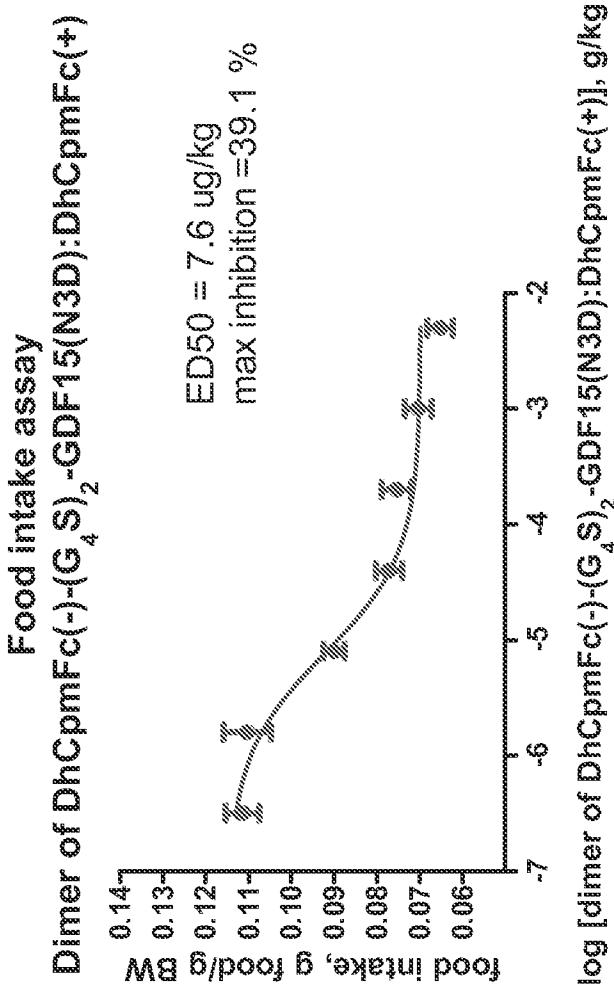
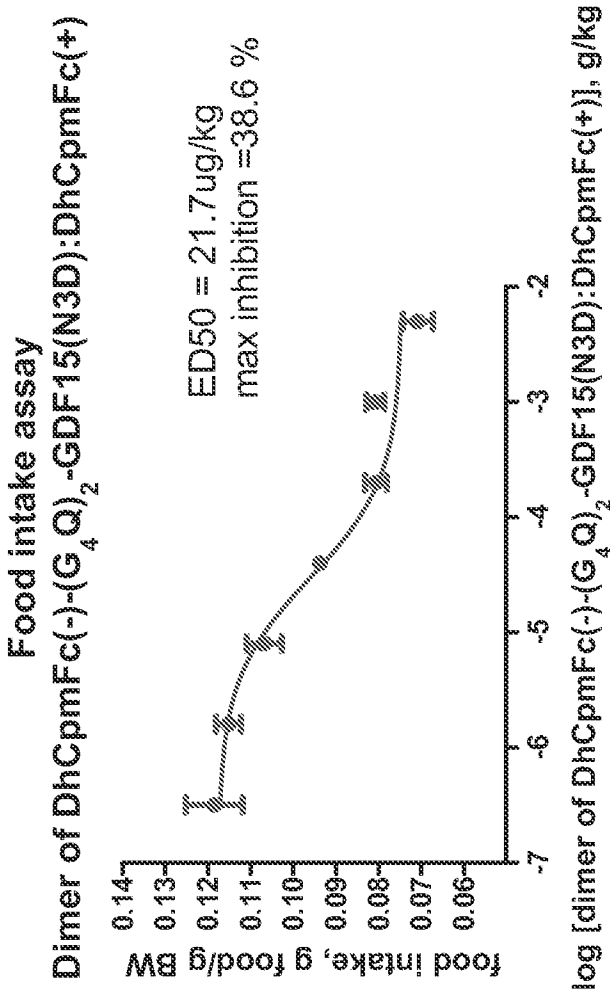


Figure 20



21/61

Figure 21

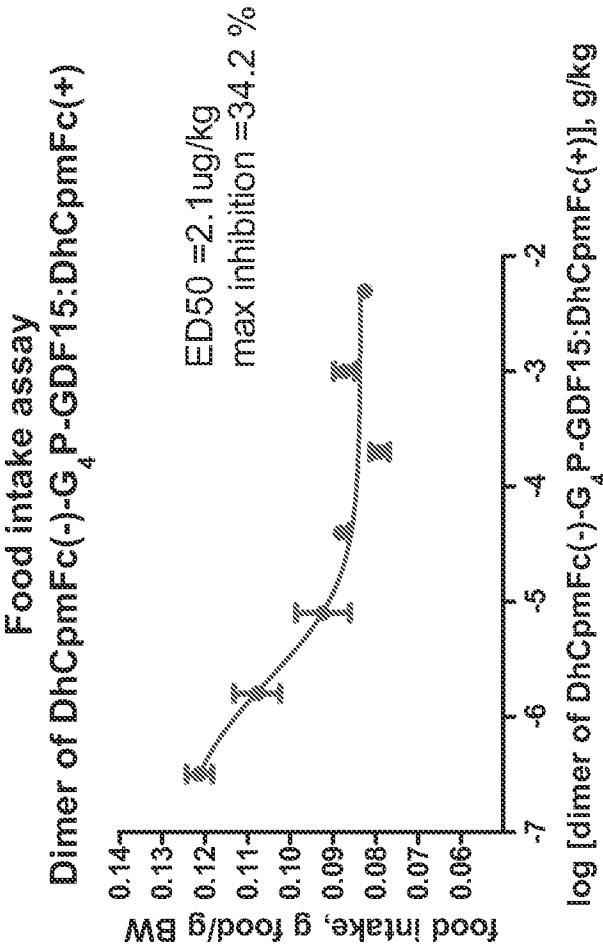
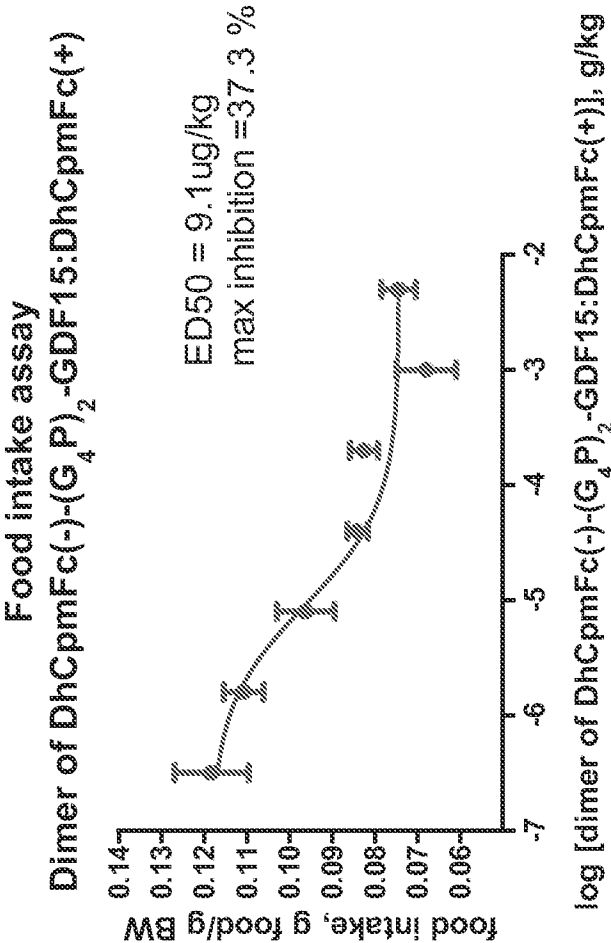


Figure 22



23/61

Figure 23

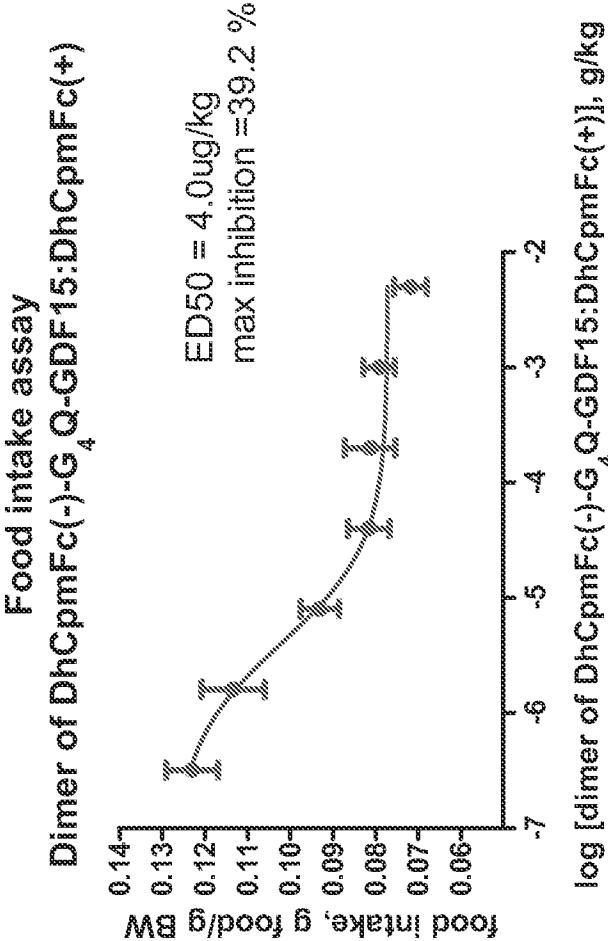


Figure 24

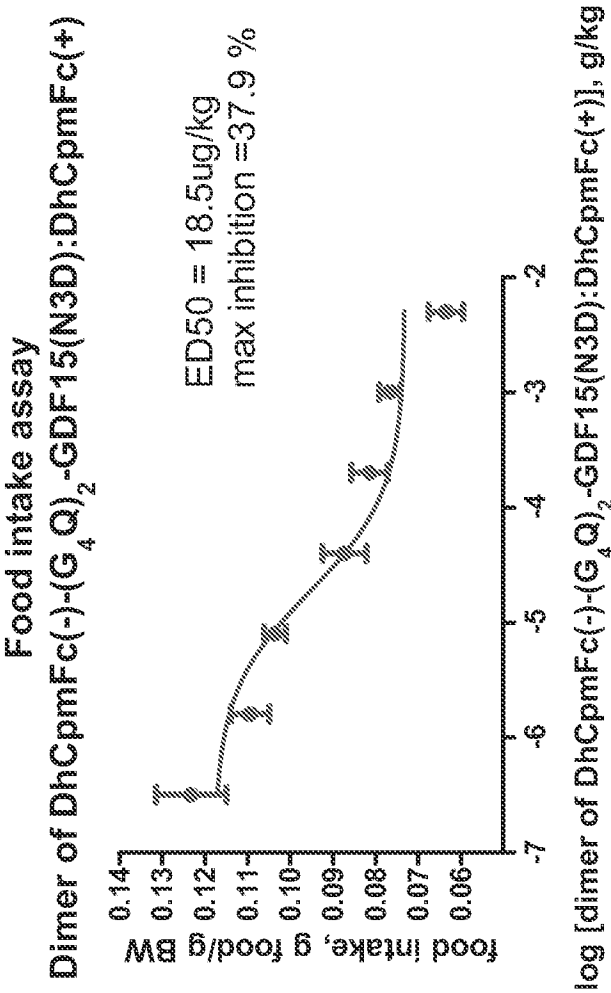


Figure 25

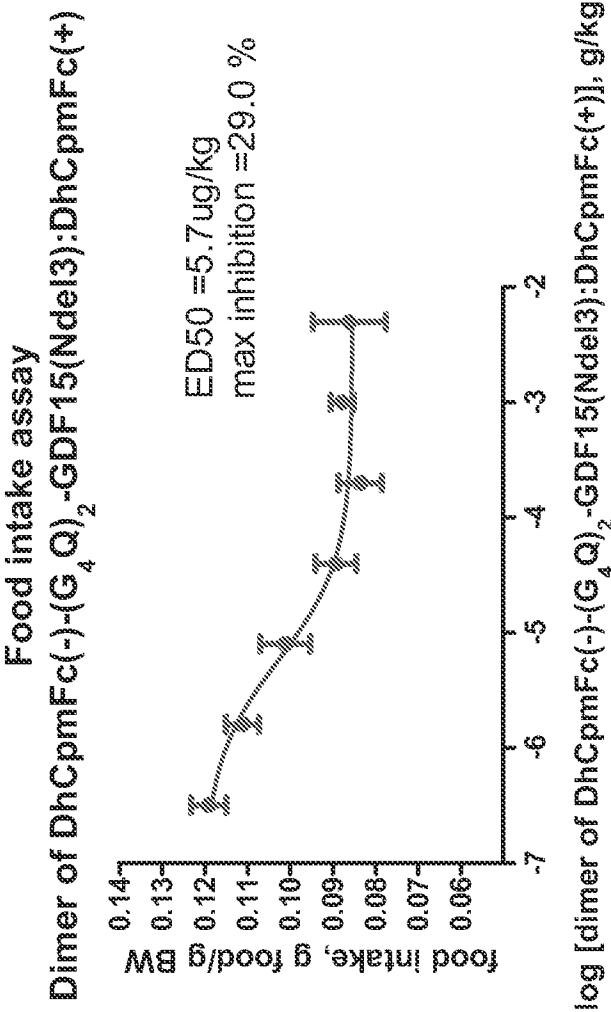
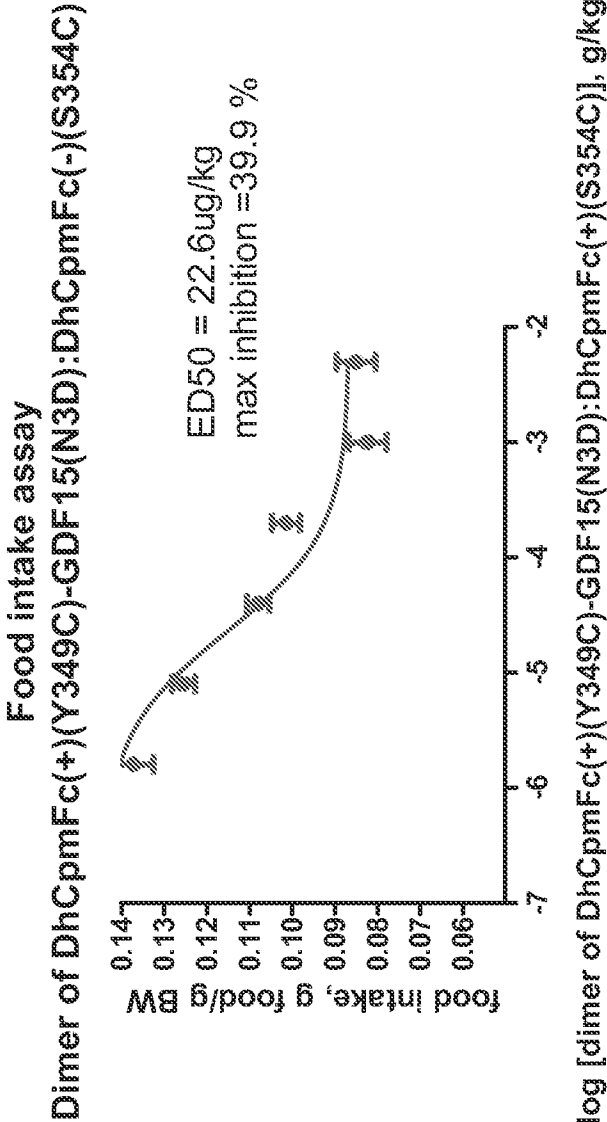
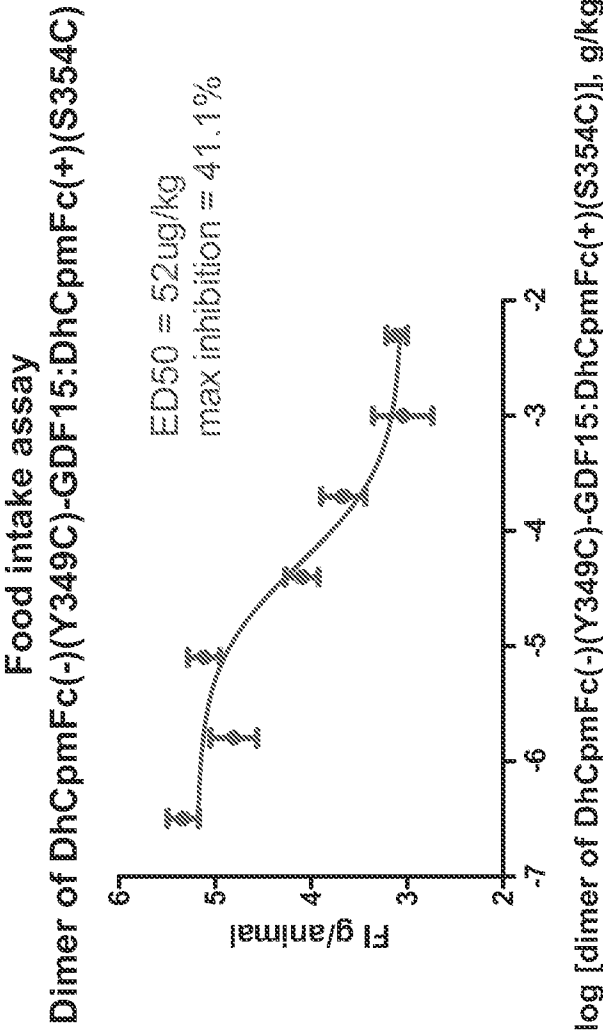


Figure 26



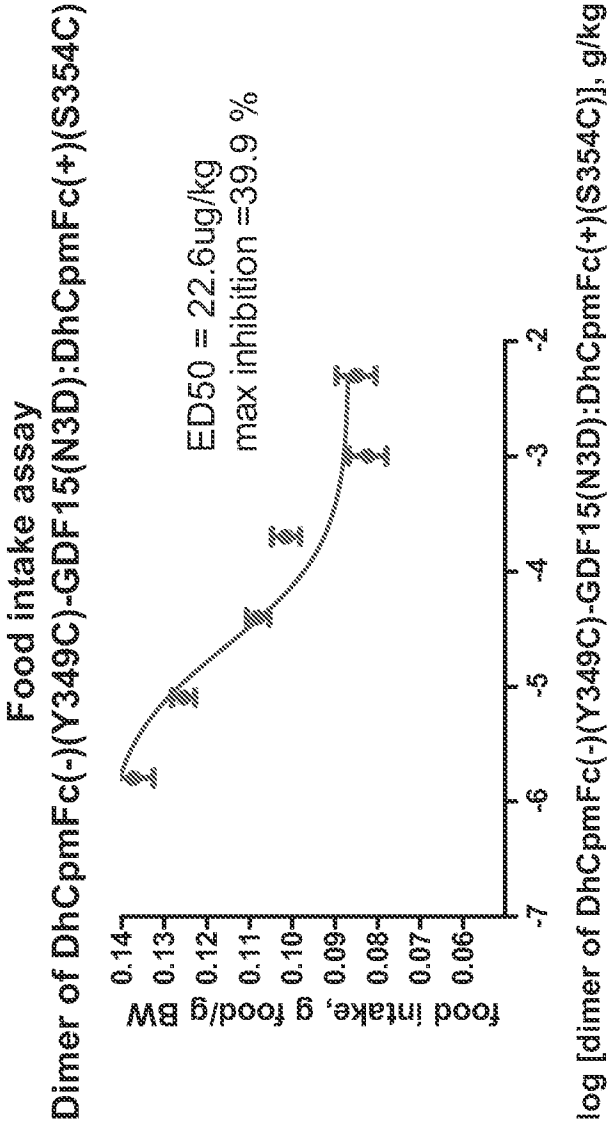
27/61

Figure 27



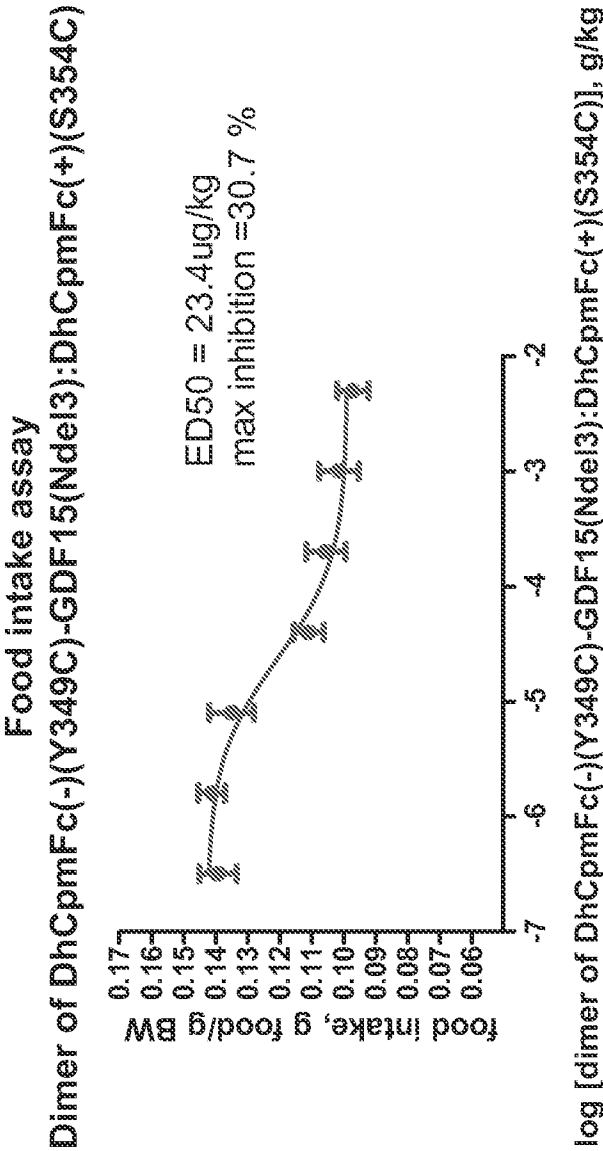
28/61

Figure 28



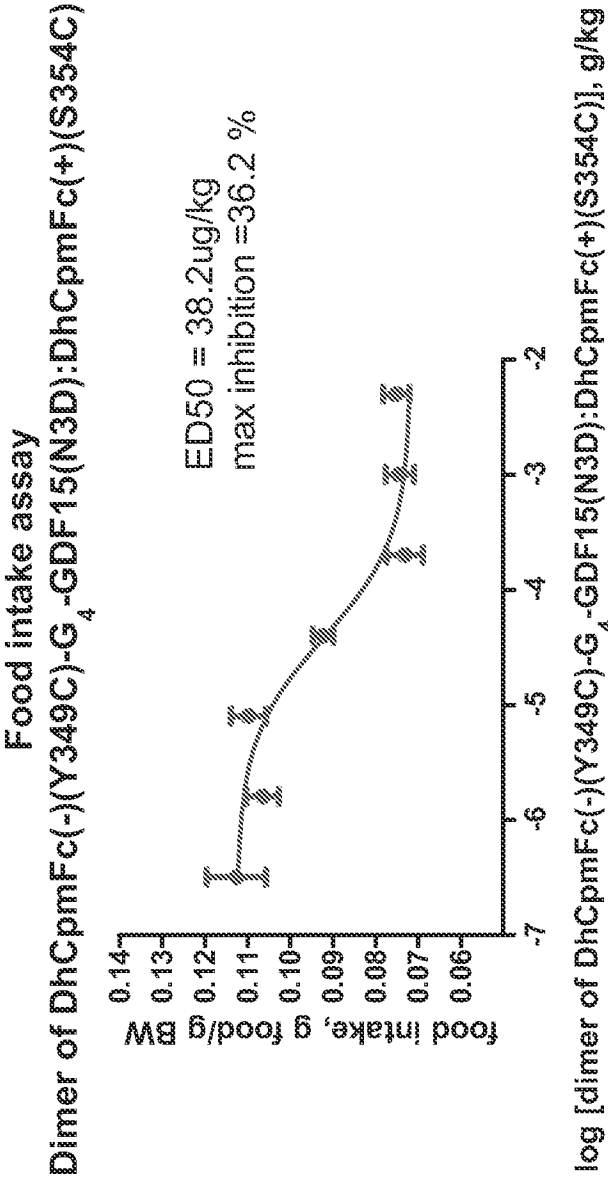
29/61

Figure 29



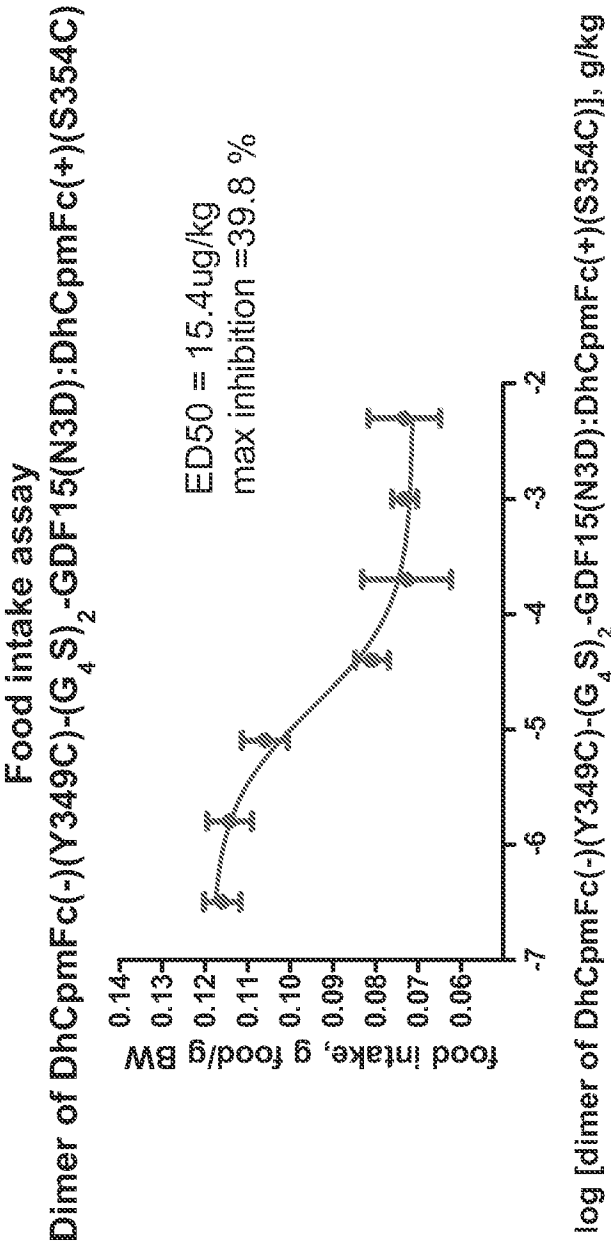
30/61

Figure 30



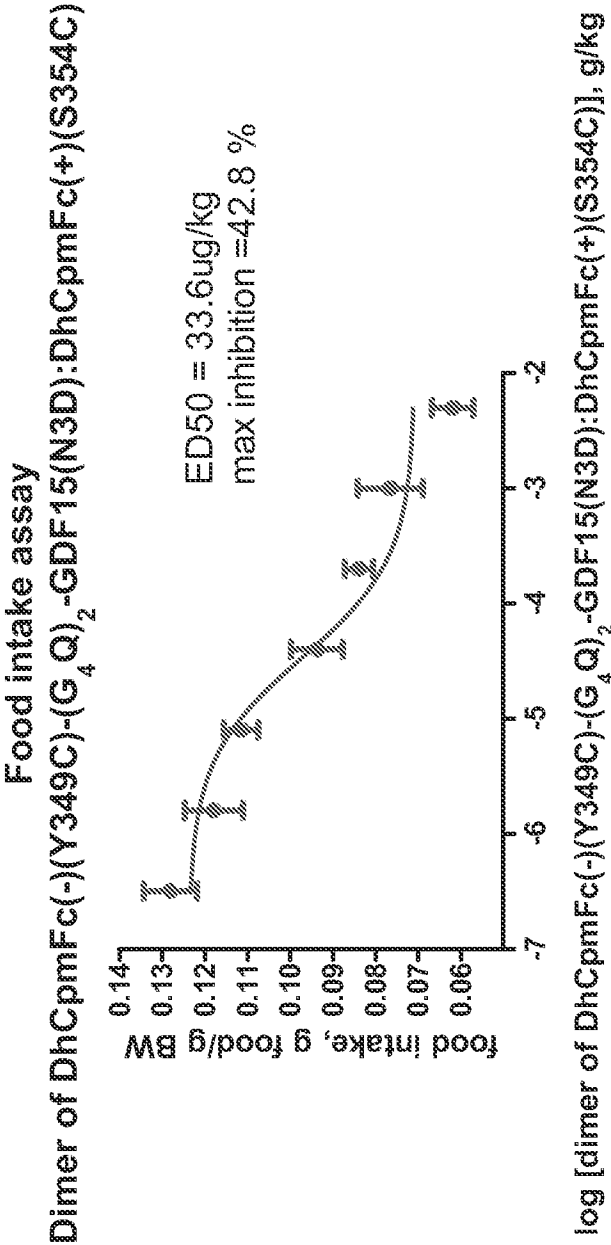
31/61

Figure 31



32/61

Figure 32



33/61

Figure 33

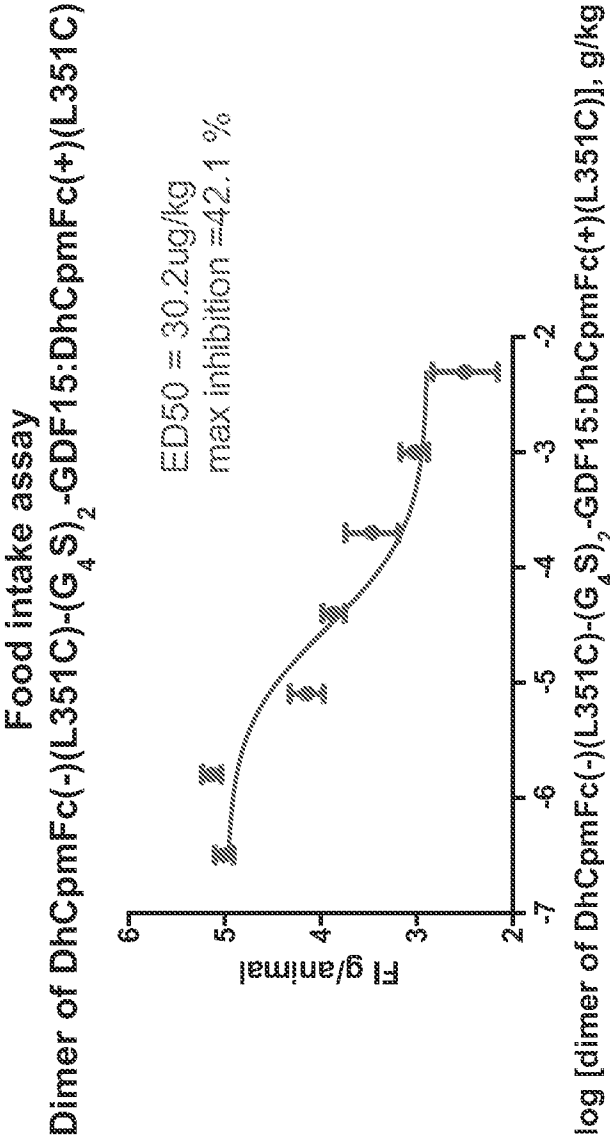


Figure 34

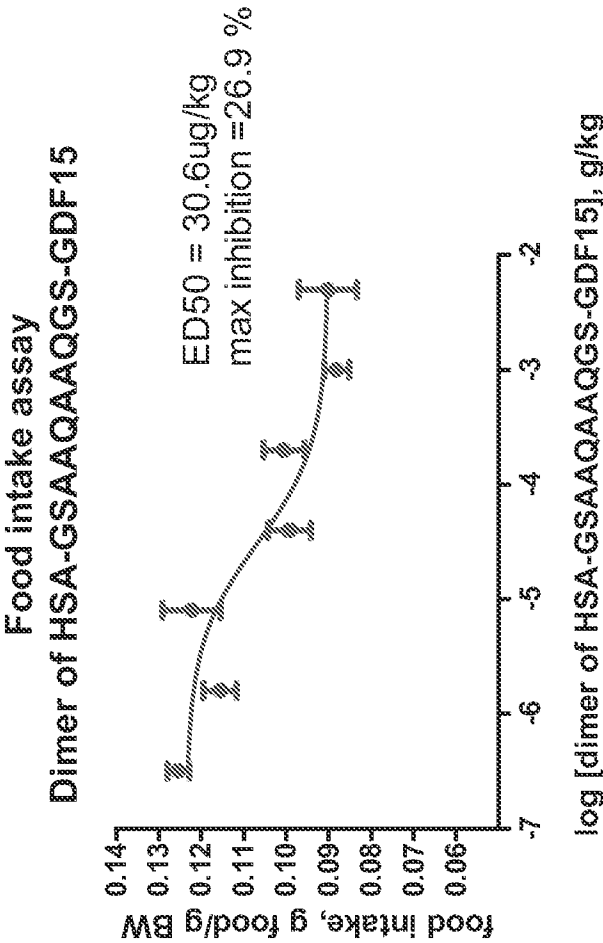
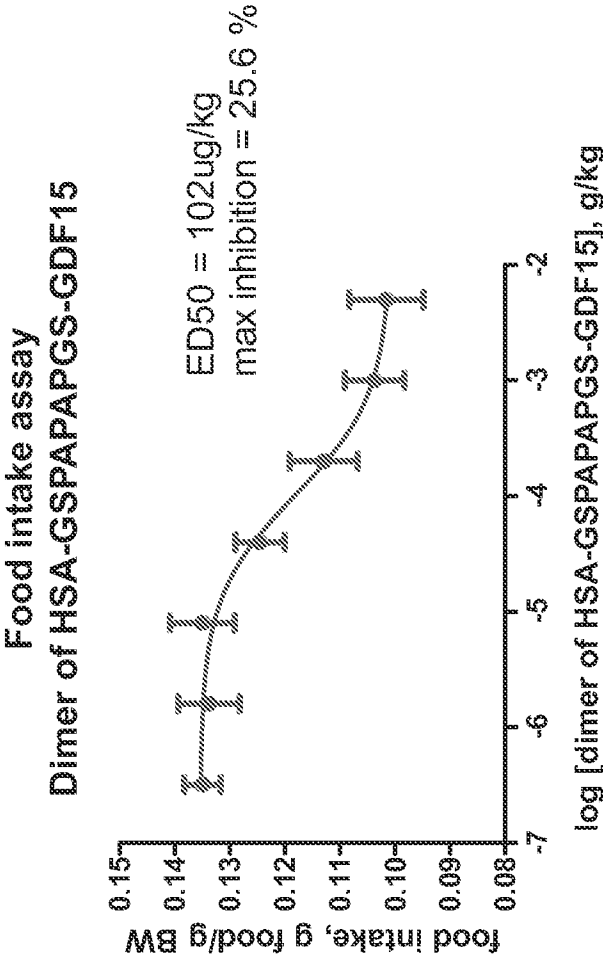
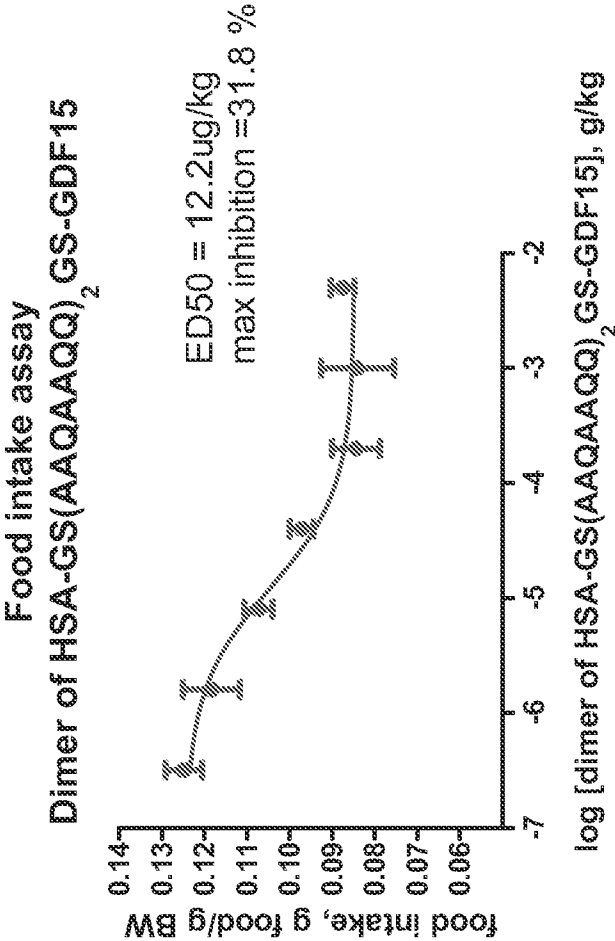


Figure 35



36/61

Figure 36



37 / 61

Figure 37

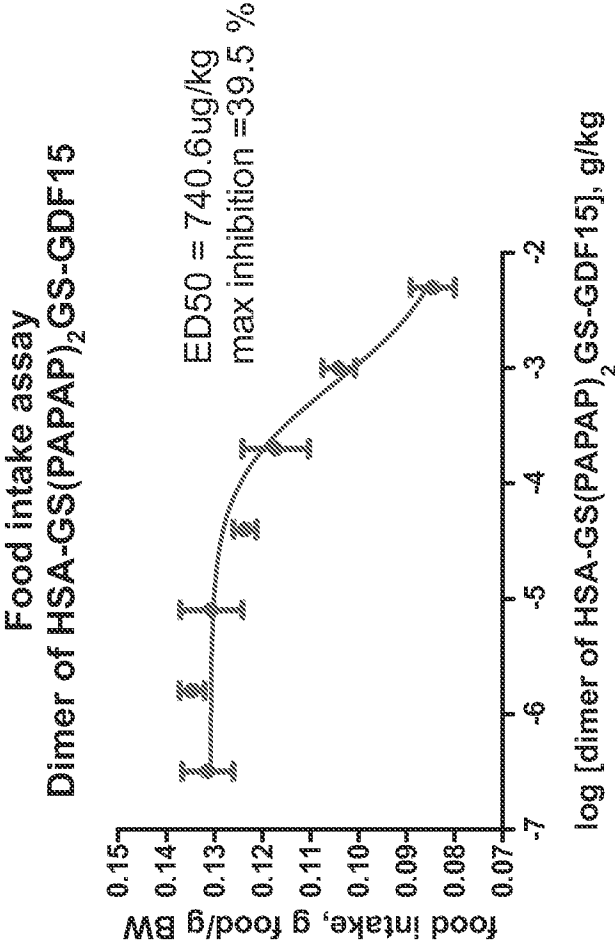


Figure 38

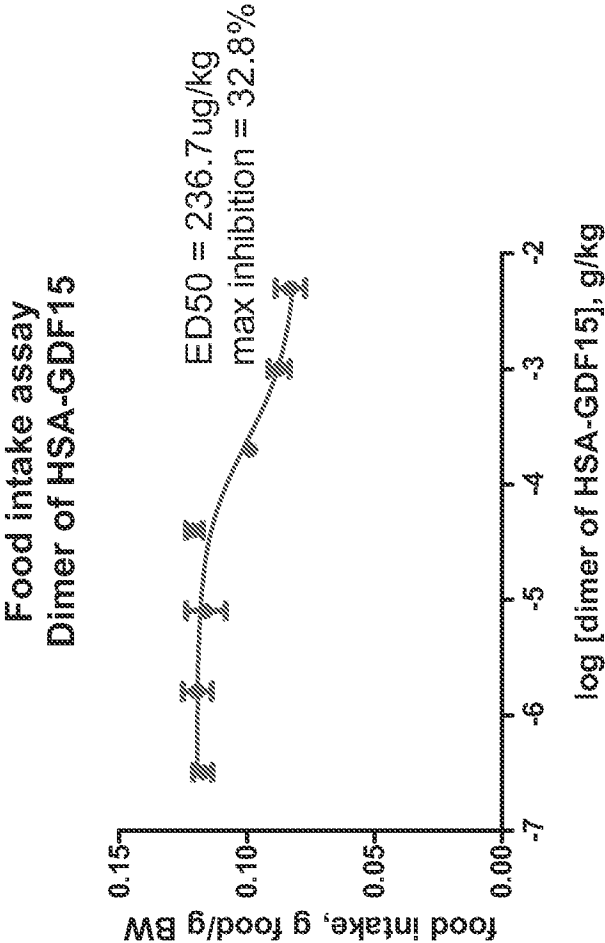
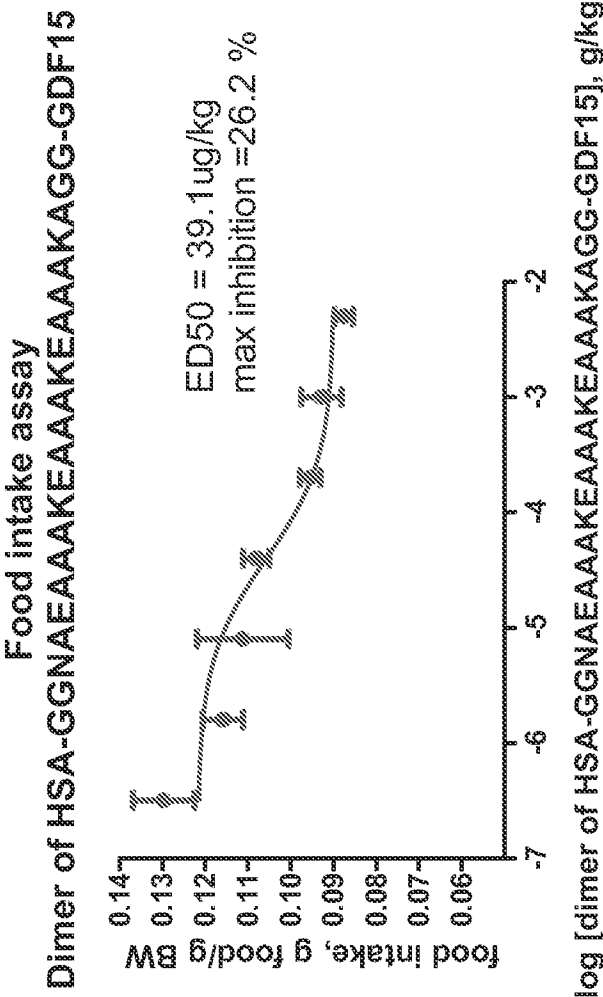


Figure 39



40/61

Figure 40

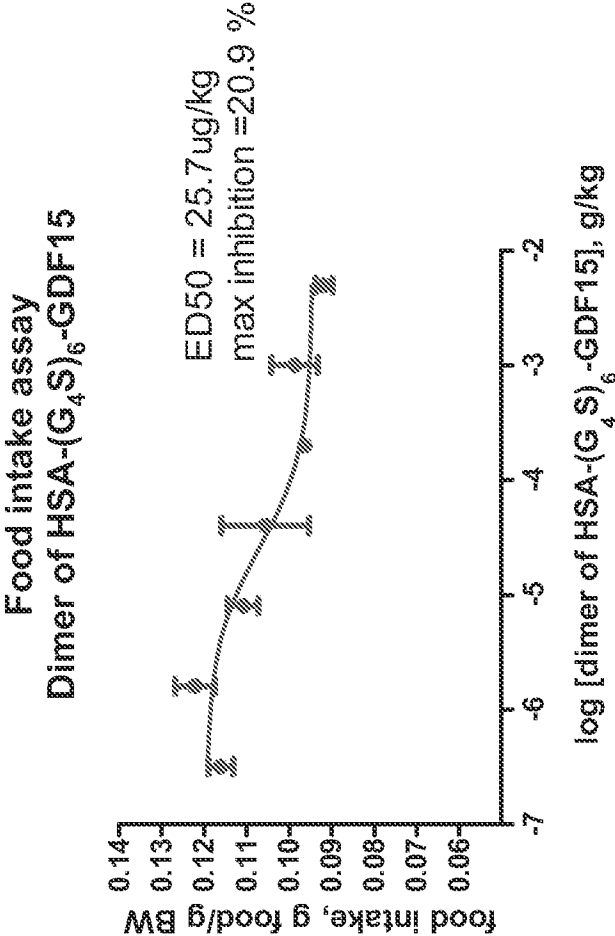
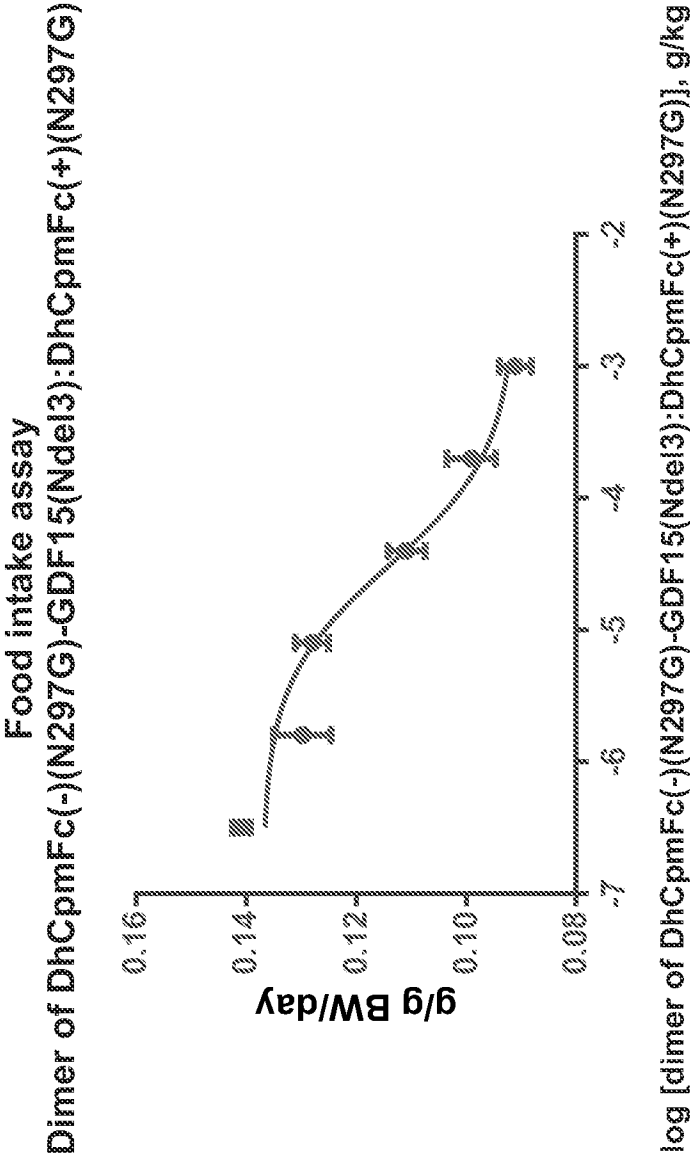
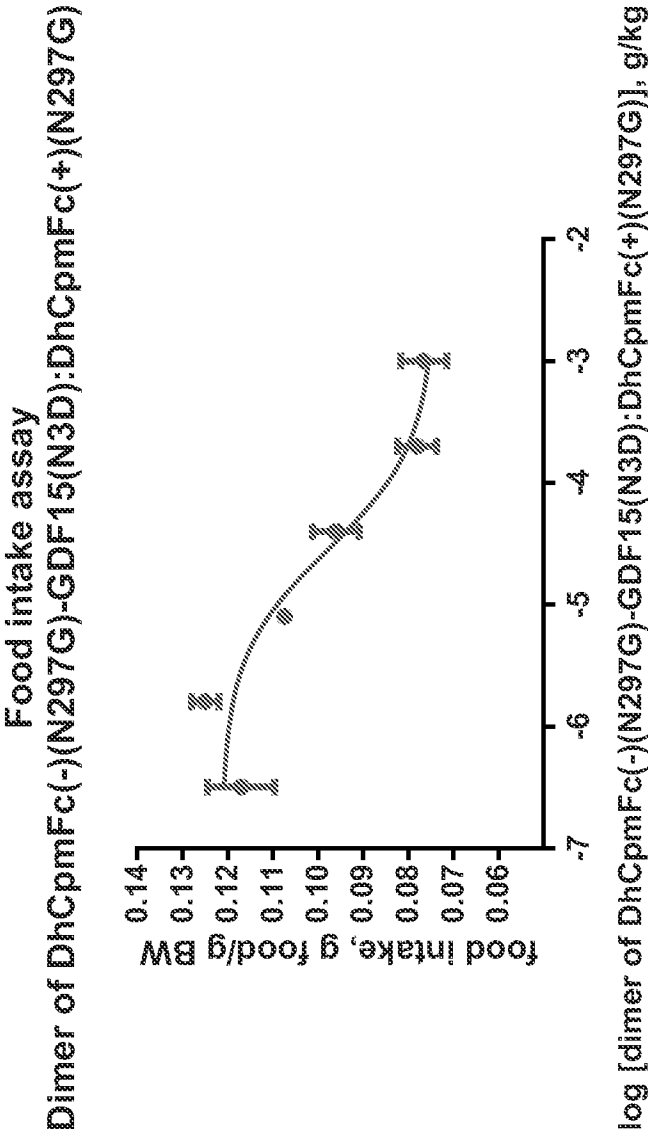


Figure 41



42/61

Figure 42



43 / 61

Figure 43

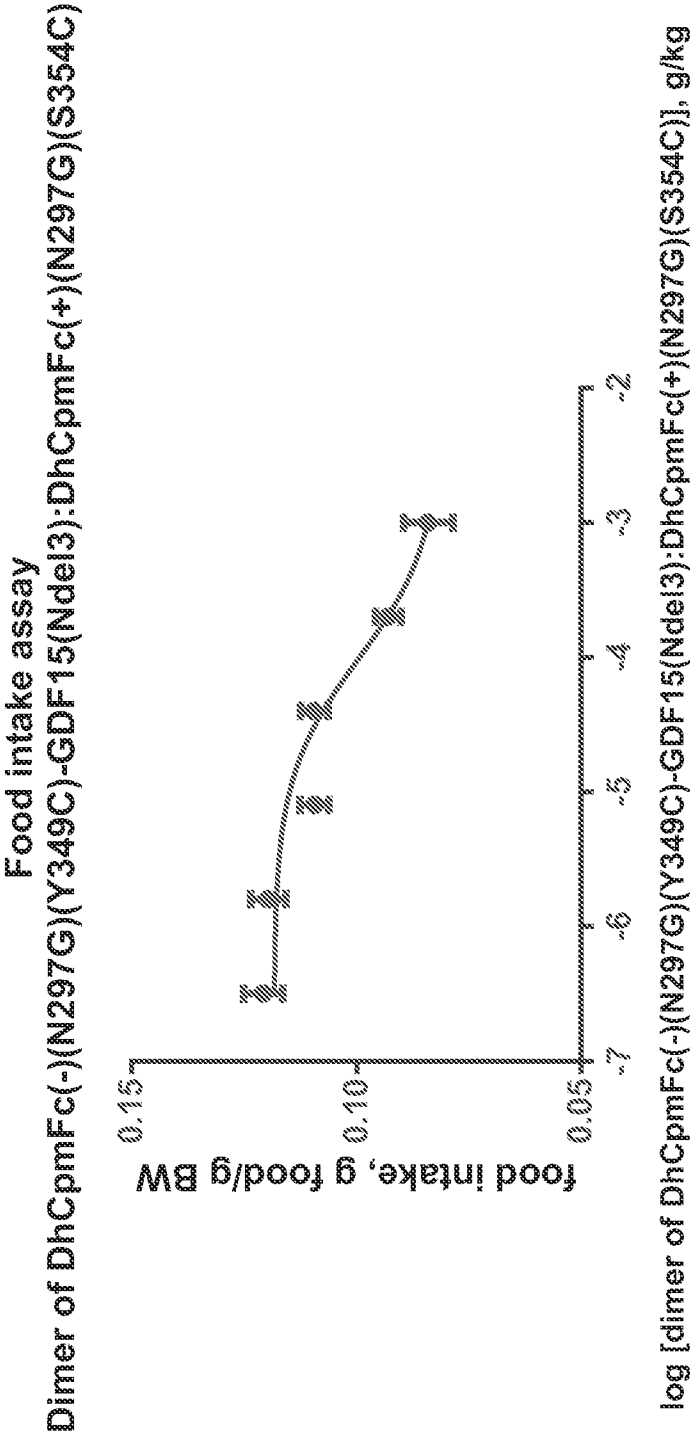
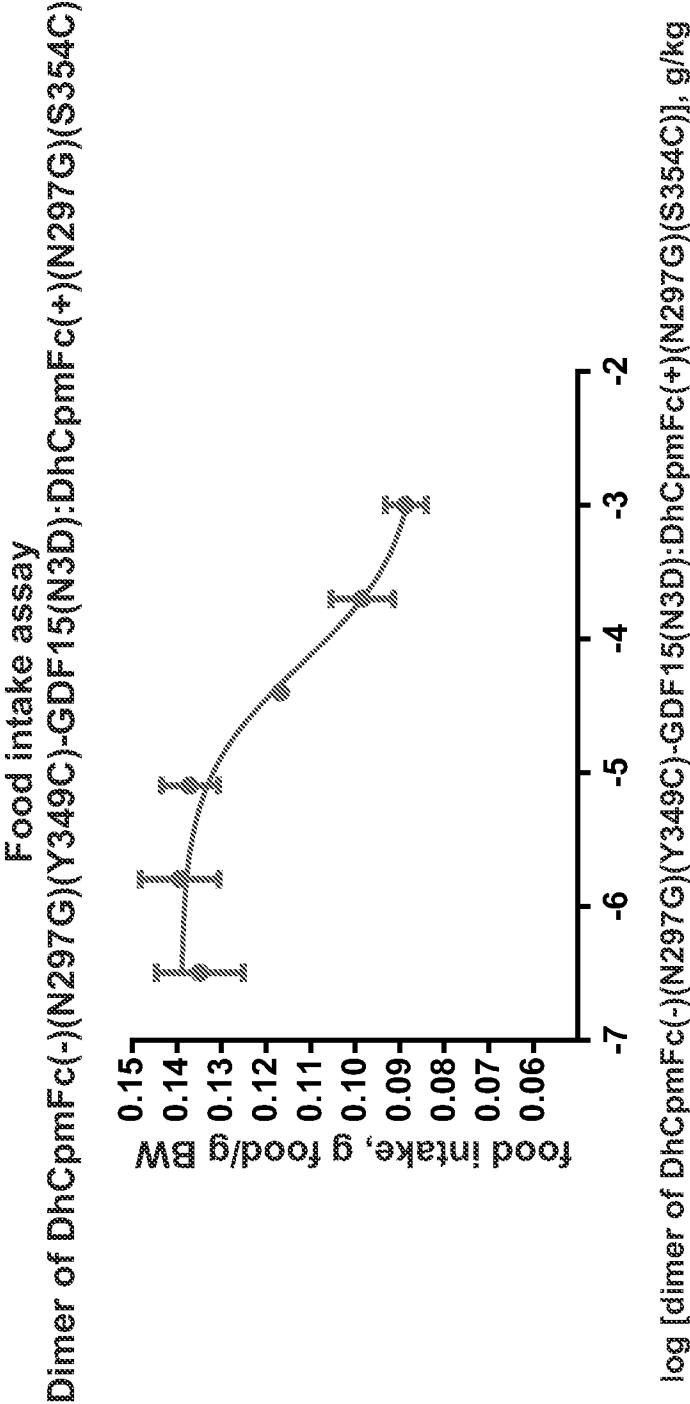


Figure 44



45/61

Figure 45

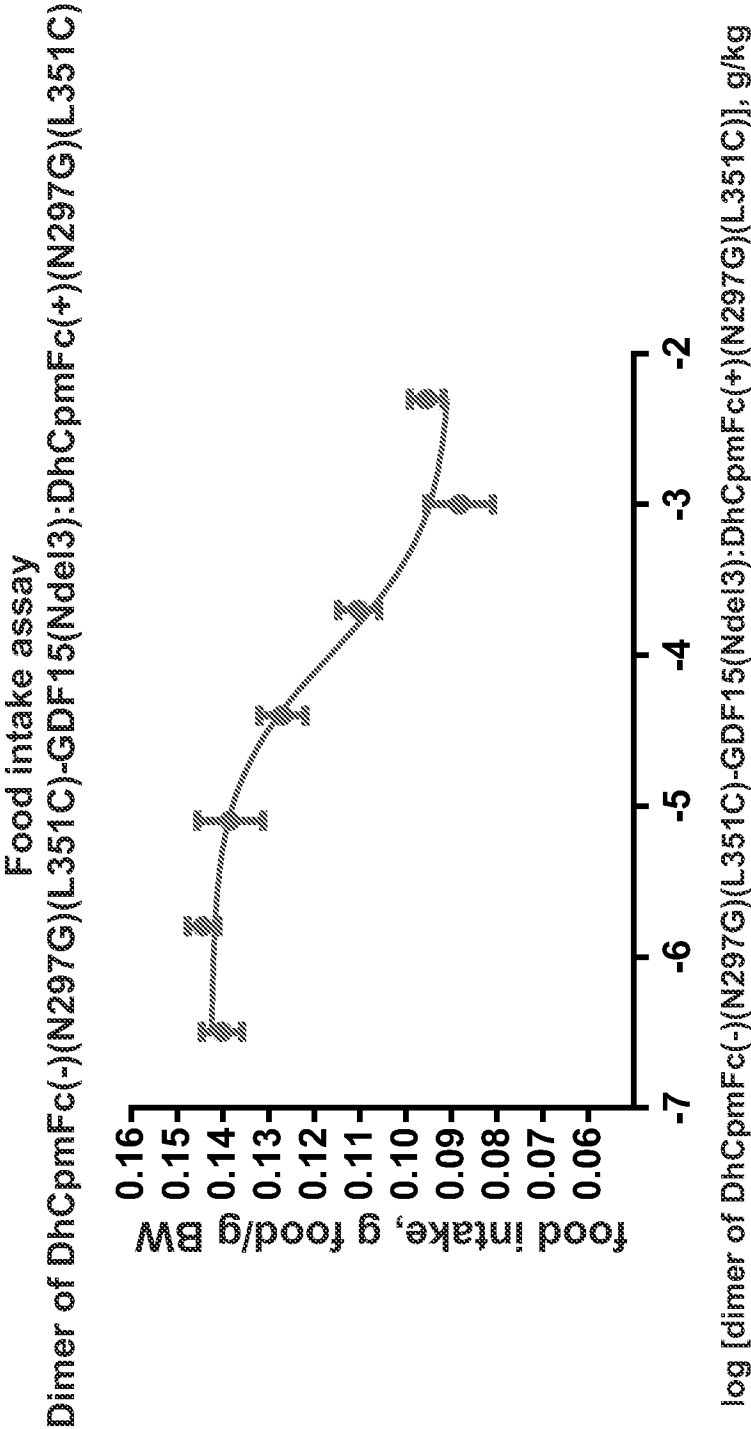
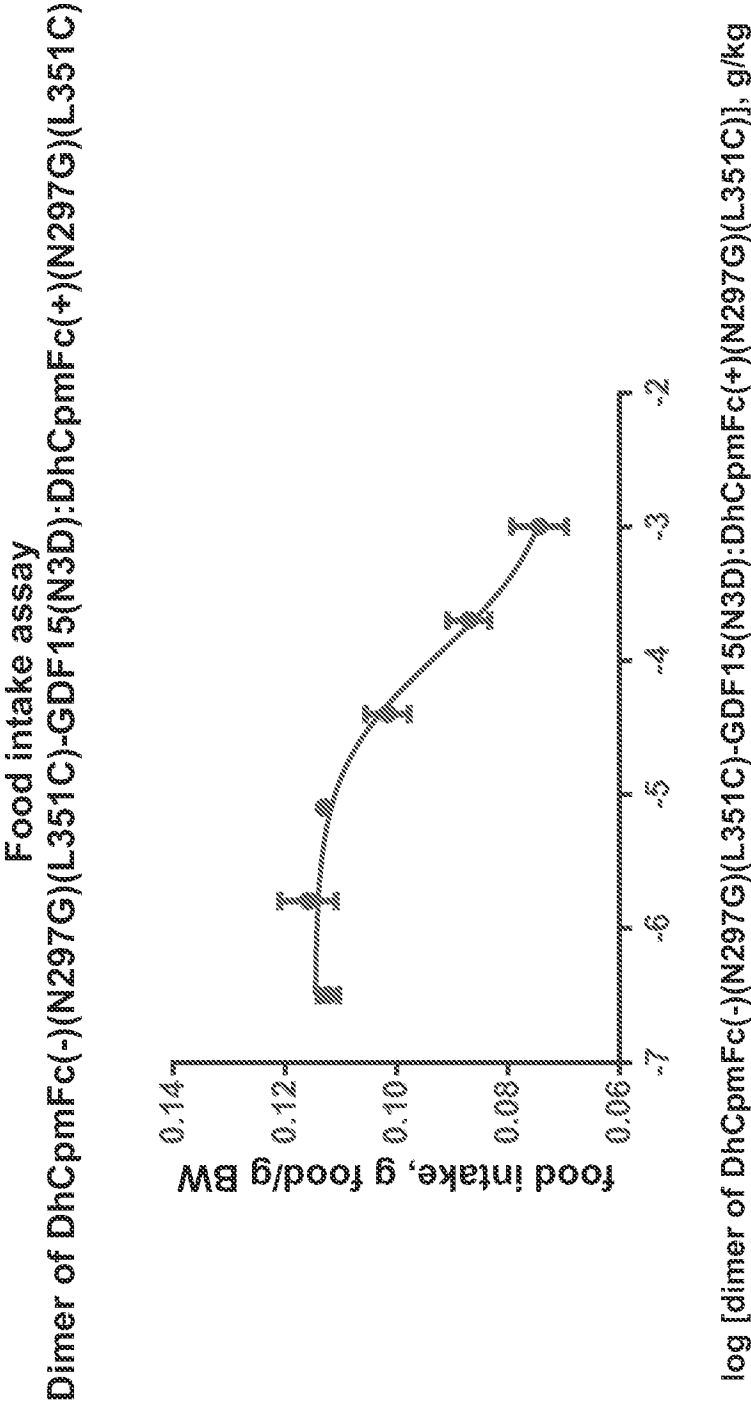
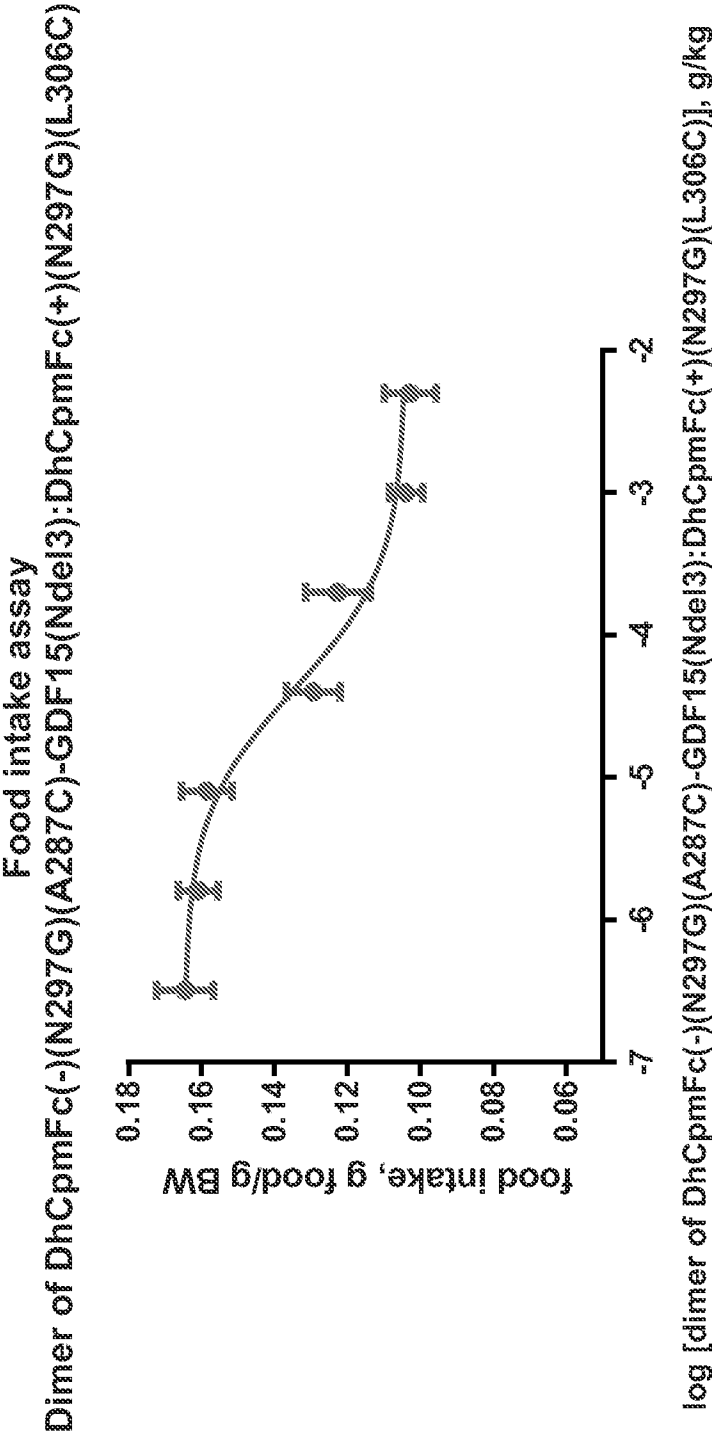


Figure 46



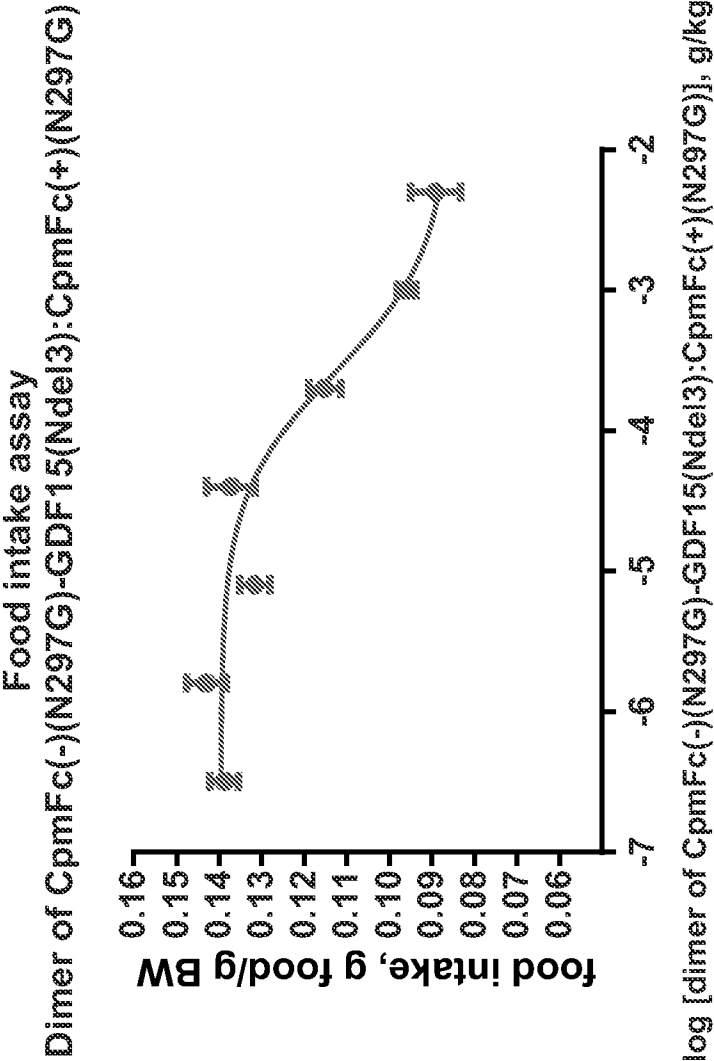
47/61

Figure 47



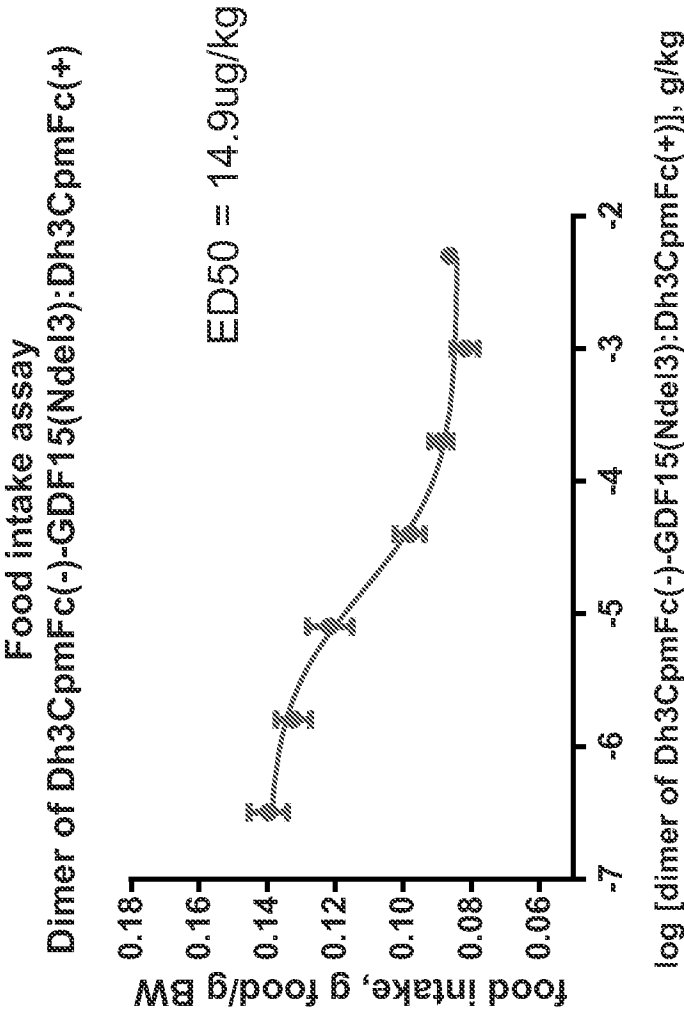
48 / 61

Figure 48



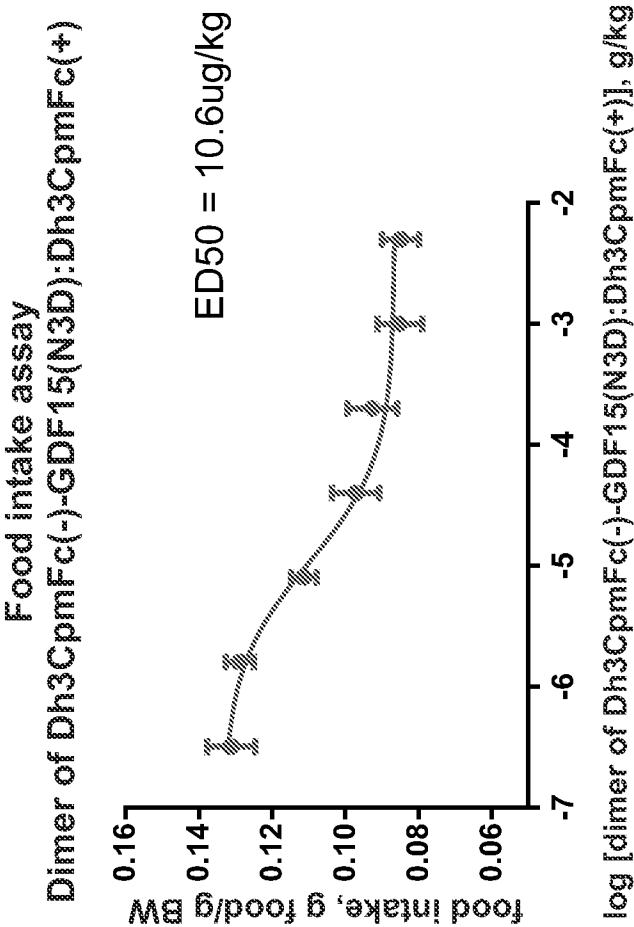
49/61

Figure 49



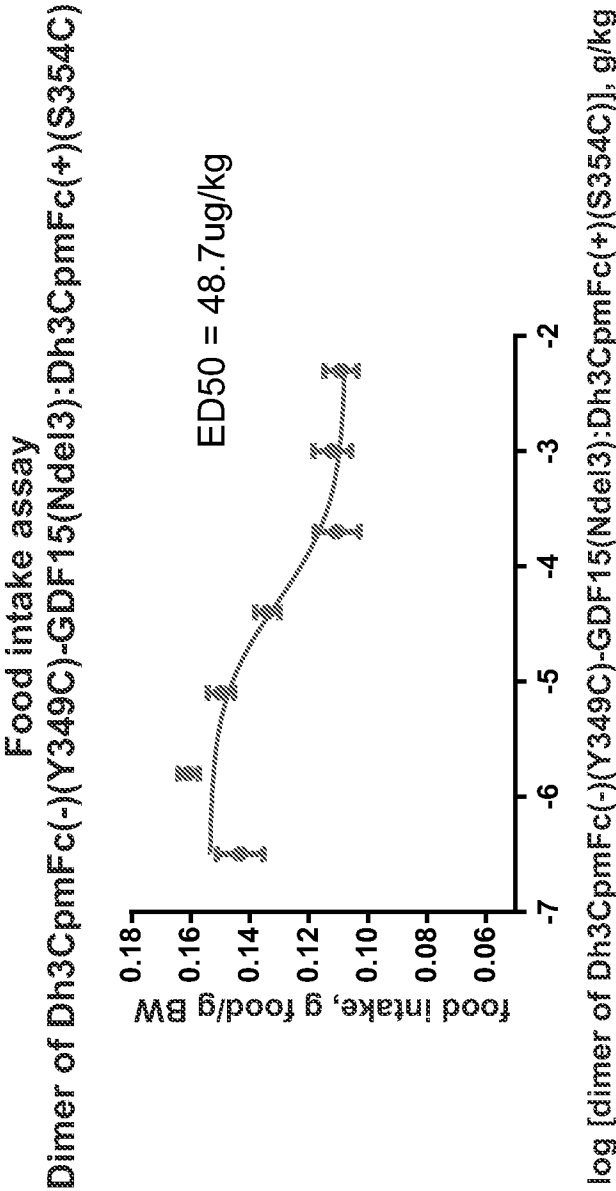
50/61

Figure 50



51/61

Figure 51



52/61

Figure 52

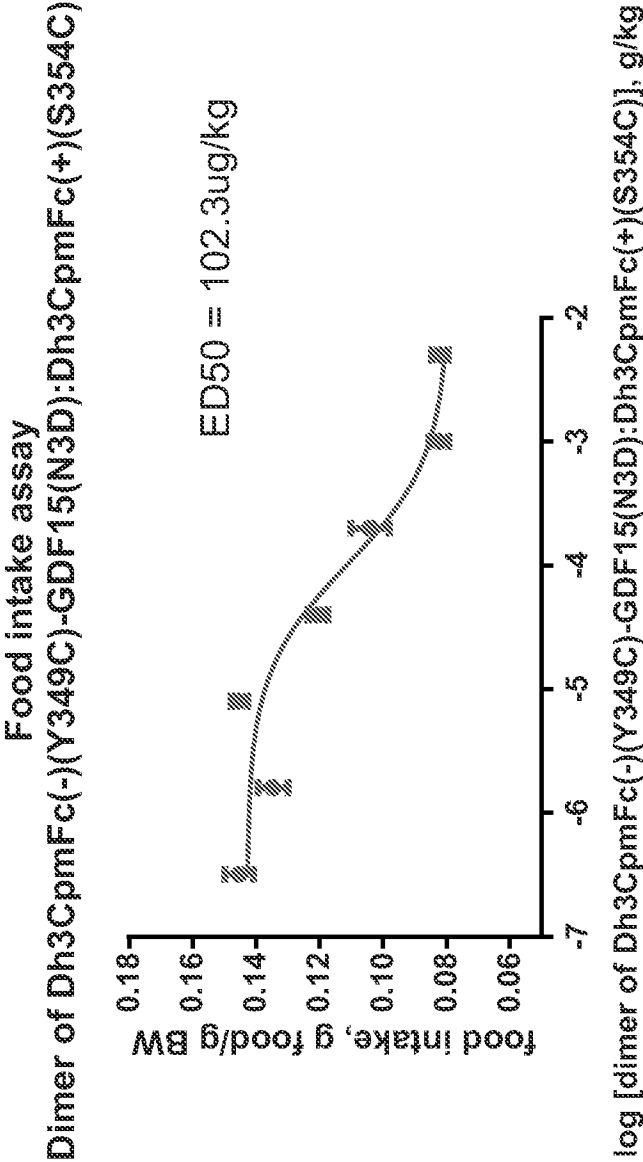
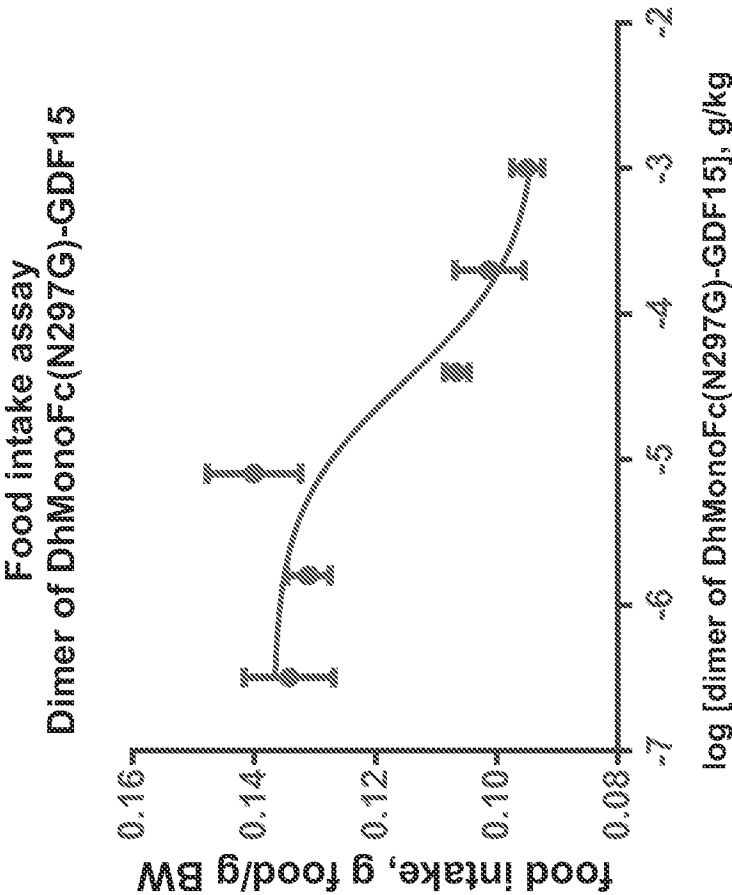
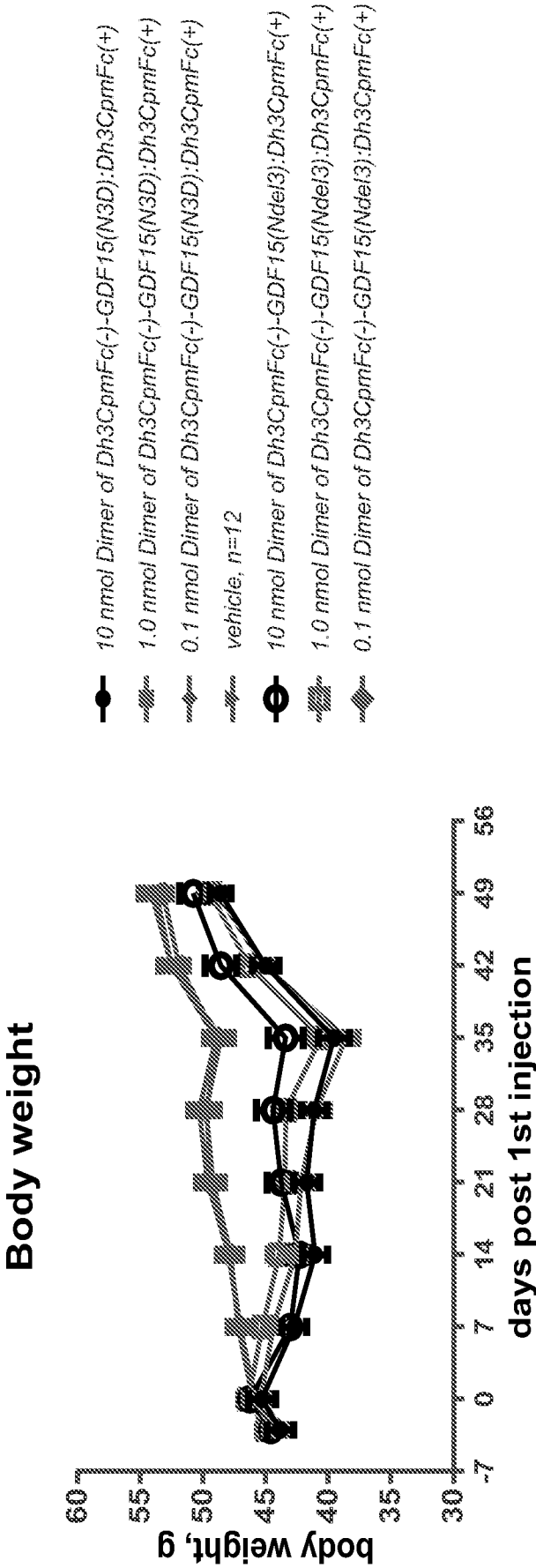


Figure 53



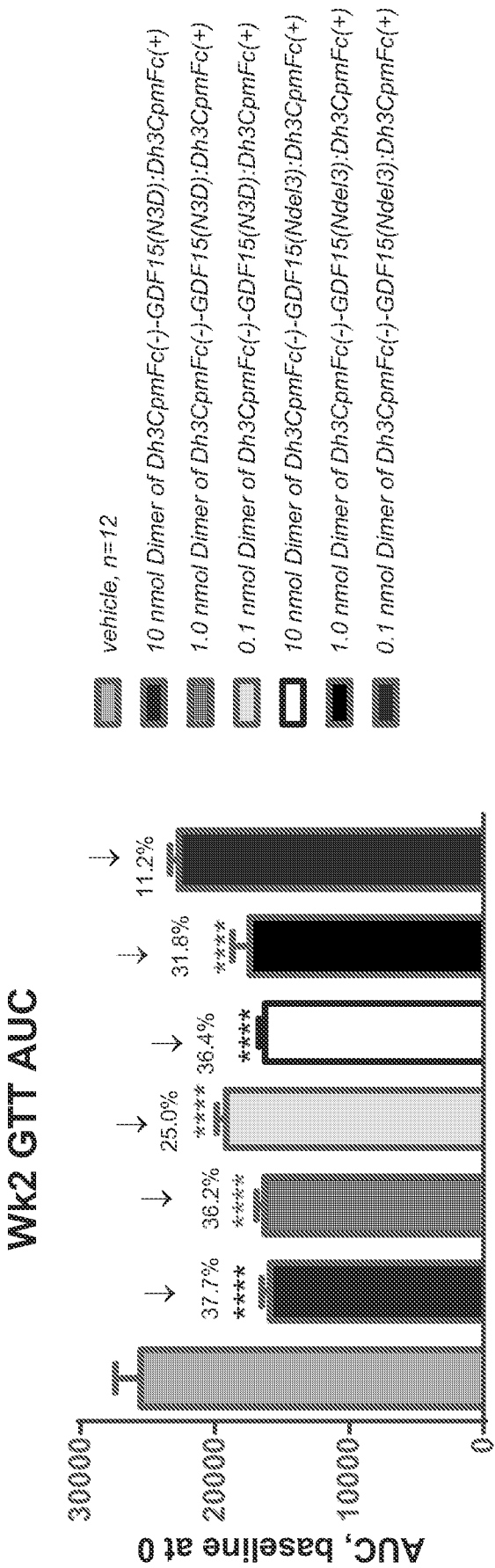
54 / 6.1

Figure 54



55/61

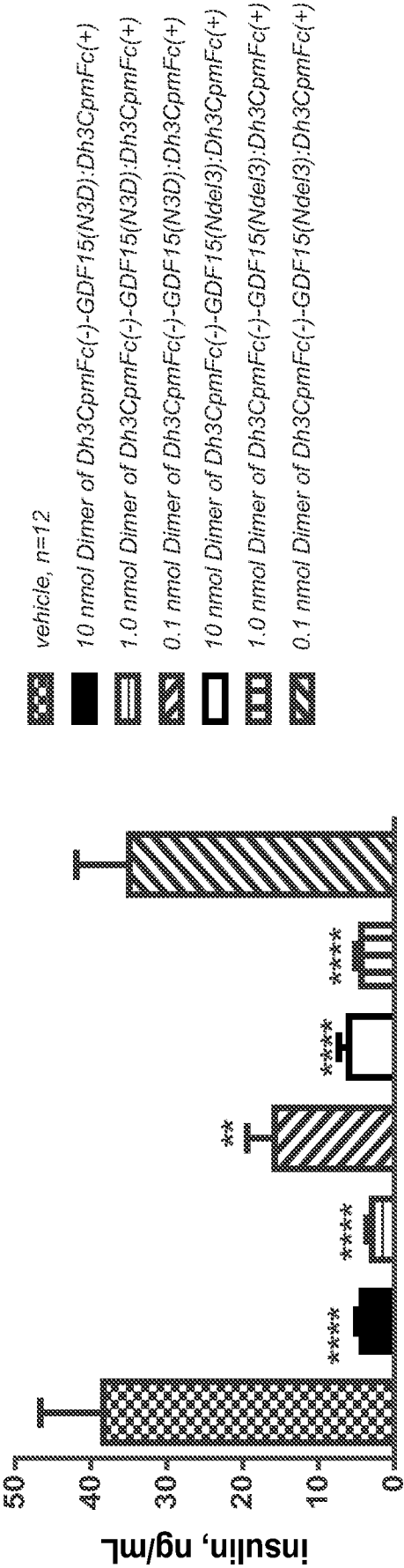
Figure 55



56/61

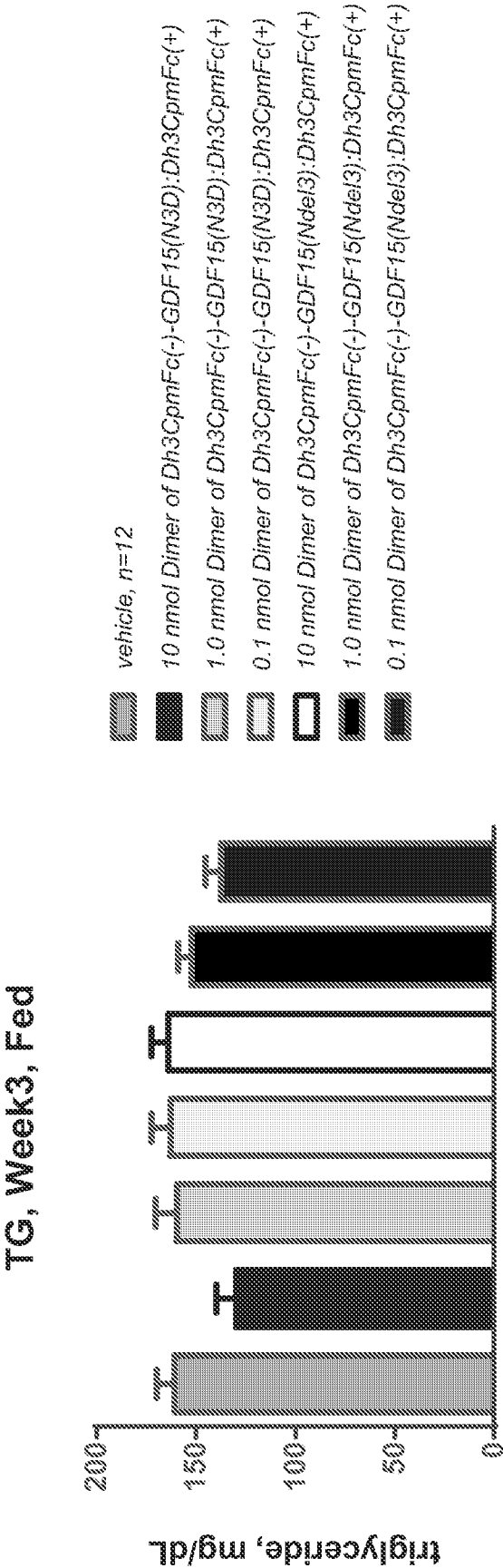
Figure 56

Insulin, Week 3, Fed



57/61

Figure 57



58/61

Figure 58

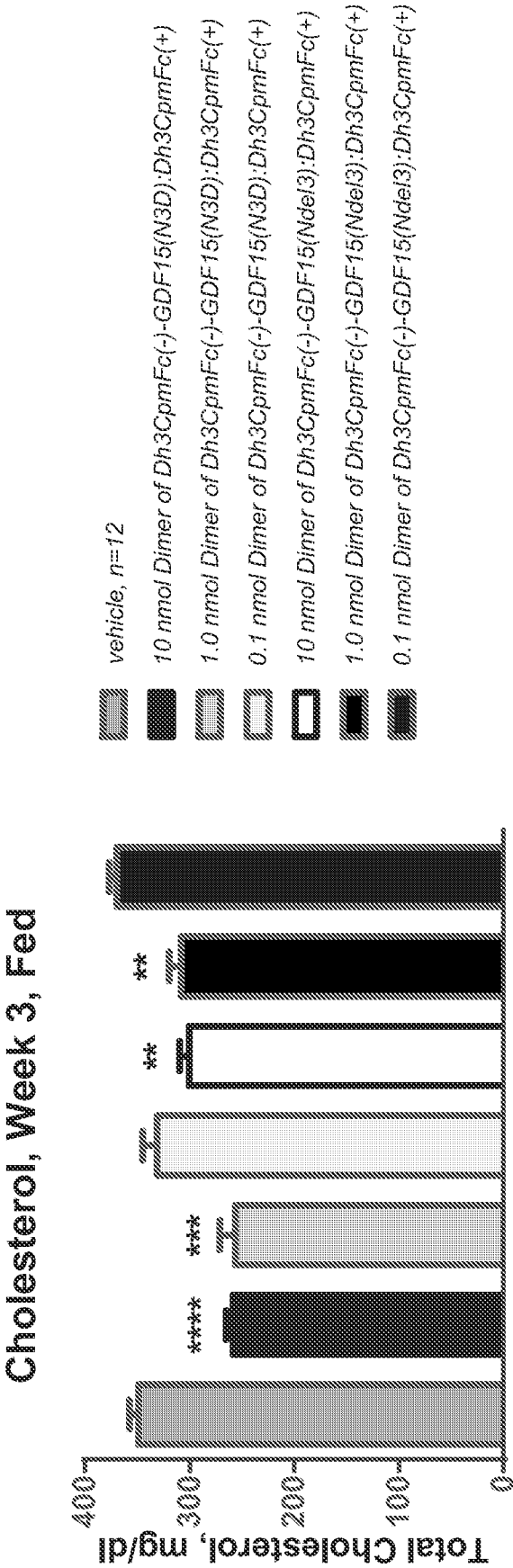
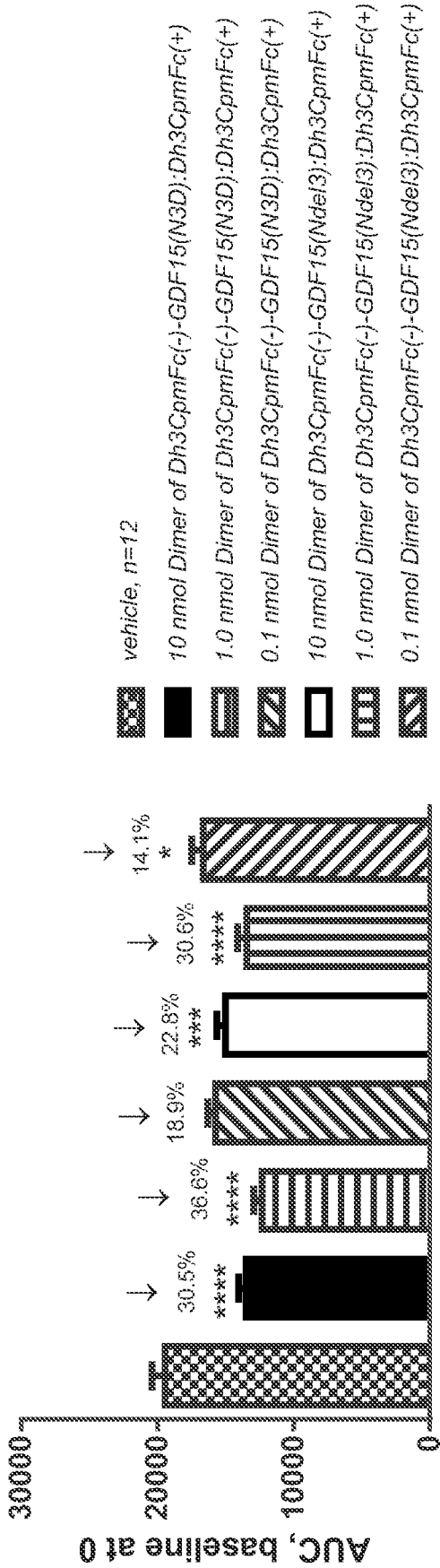


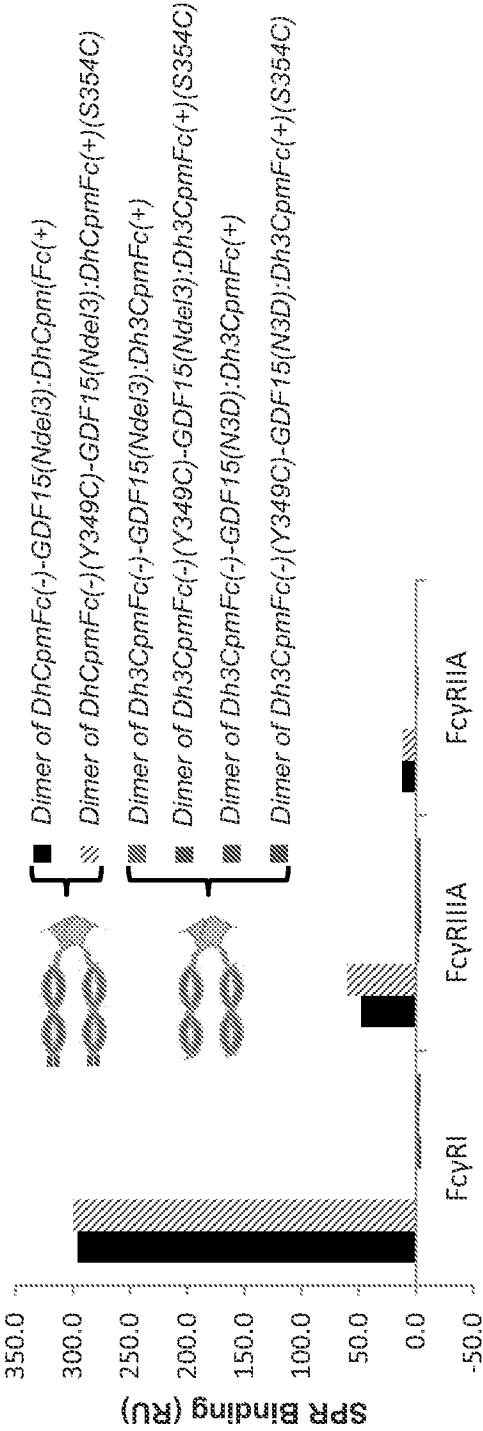
Figure 59

Wk5 GTT AUC



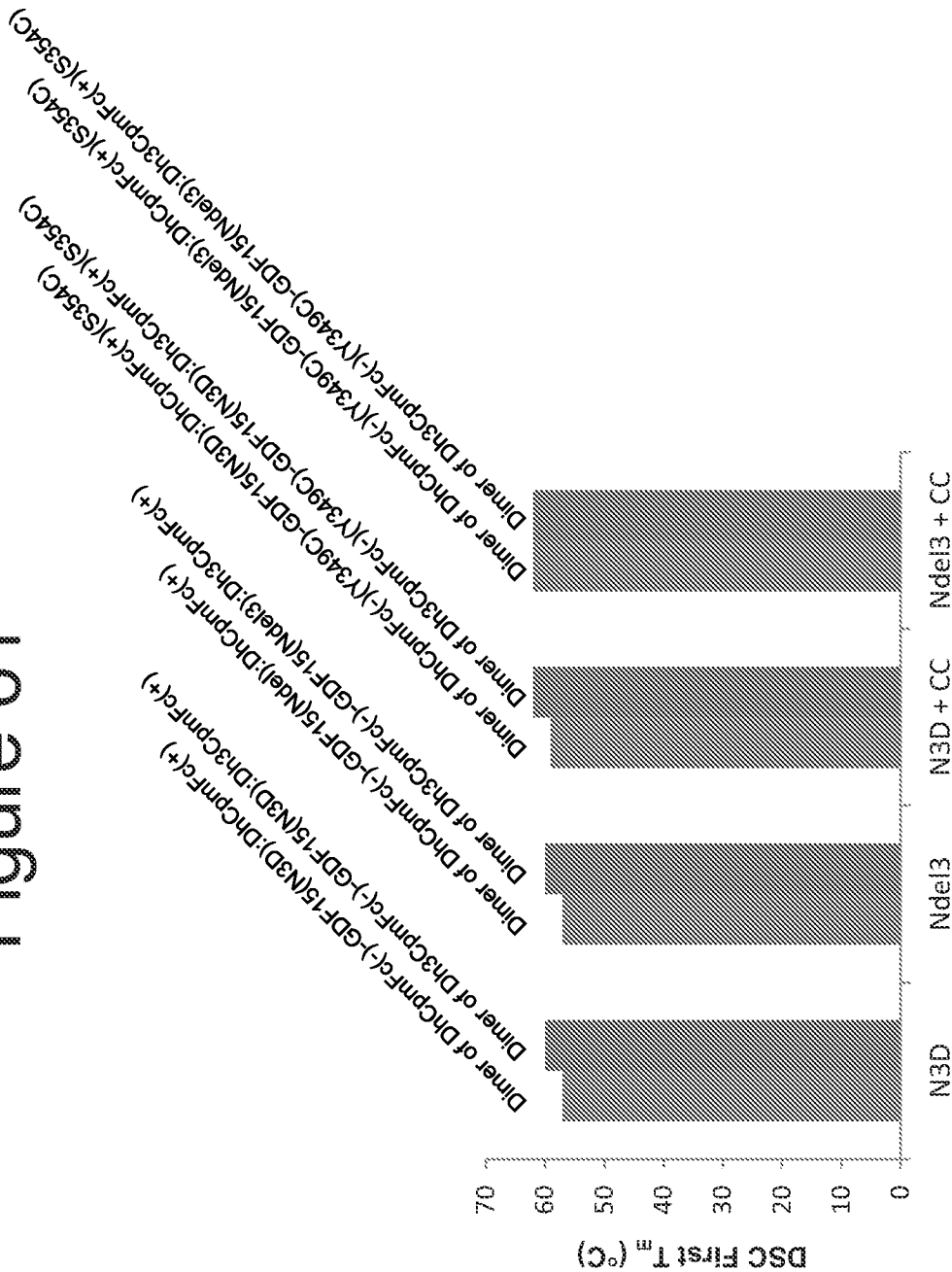
60/61

Figure 60



61/61

Figure 61



INTERNATIONAL SEARCH REPORT

International application No

PCT/US2014/049254

A. CLASSIFICATION OF SUBJECT MATTER

INV. C07K14/495 C12N15/62
ADD.

According to International Patent Classification (IPC) or to both national classification and IPC

B. FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols)

C07K C12N

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the international search (name of data base and, where practicable, search terms used)

EPO-Internal, BIOSIS, EMBASE, WPI Data

C. DOCUMENTS CONSIDERED TO BE RELEVANT

Category*	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
Y	WO 99/06445 A1 (UNIV JOHNS HOPKINS MED [US]; LEE SE JIN [US]; HUYNH THANH V [US]; SEBA) 11 February 1999 (1999-02-11) abstract figure 1	1-25
Y	WO 2010/048670 A1 (ST VINCENTS HOSP SYDNEY [AU]; BREIT SAMUEL NORBERT [AU]; BROWN DAVID A) 6 May 2010 (2010-05-06) abstract page 1, line 16 - line 29	1-25
Y	WO 2006/000448 A2 (MERCK PATENT GMBH [DE]; GILLIES STEPHEN D [US]; WATKINS NIGEL JOHN [GB]) 5 January 2006 (2006-01-05) abstract page 3, line 7 - line 14	1-25
	----- -/-	



Further documents are listed in the continuation of Box C.



See patent family annex.

* Special categories of cited documents :

"A" document defining the general state of the art which is not considered to be of particular relevance

"E" earlier application or patent but published on or after the international filing date

"L" document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified)

"O" document referring to an oral disclosure, use, exhibition or other means

"P" document published prior to the international filing date but later than the priority date claimed

"T" later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention

"X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone

"Y" document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art

"&" document member of the same patent family

Date of the actual completion of the international search

13 November 2014

Date of mailing of the international search report

10/12/2014

Name and mailing address of the ISA/

European Patent Office, P.B. 5818 Patentlaan 2
NL - 2280 HV Rijswijk
Tel. (+31-70) 340-2040,
Fax: (+31-70) 340-3016

Authorized officer

Grötzinger, Thilo

INTERNATIONAL SEARCH REPORT

International application No

PCT/US2014/049254

C(Continuation). DOCUMENTS CONSIDERED TO BE RELEVANT

Category*	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
Y	WO 2011/064758 A2 (PFIZER LTD [GB]; CIARAMELLA GIUSEPPE [DE]; FLORES MARIA VICTORIA [GB];) 3 June 2011 (2011-06-03) abstract page 4, line 1 - line 21 -----	1-25
X,P	WO 2013/148117 A1 (NGM BIOPHARMACEUTICALS INC [US]) 3 October 2013 (2013-10-03) abstract page 2, line 21 - line 33 page 3, line 22 - line 25 -----	1-25
X,P	WO 2013/113008 A1 (AMGEN INC [US]) 1 August 2013 (2013-08-01) abstract page 2, line 28 - line 30 page 3, line 7 - line 8 -----	1-25

INTERNATIONAL SEARCH REPORT

Information on patent family members

International application No

PCT/US2014/049254

Patent document cited in search report		Publication date	Patent family member(s)		Publication date
WO 9906445	A1	11-02-1999	AU	8591198 A	22-02-1999
			US	6420543 B1	16-07-2002
			US	2003023073 A1	30-01-2003
			WO	9906445 A1	11-02-1999

WO 2010048670	A1	06-05-2010	CA	2740632 A1	06-05-2010
			CN	102203619 A	28-09-2011
			EP	2350669 A1	03-08-2011
			ES	2434996 T3	18-12-2013
			HK	1154652 A1	25-04-2014
			JP	5444363 B2	19-03-2014
			JP	2012507012 A	22-03-2012
			US	2012107420 A1	03-05-2012
			US	2014205684 A1	24-07-2014
			WO	2010048670 A1	06-05-2010

WO 2006000448	A2	05-01-2006	AT	443720 T	15-10-2009
			AU	2005256519 A1	05-01-2006
			CA	2572015 A1	05-01-2006
			EP	1761559 A2	14-03-2007
			ES	2332924 T3	15-02-2010
			JP	4808709 B2	02-11-2011
			JP	2008508862 A	27-03-2008
			US	2006228332 A1	12-10-2006
			US	2009191154 A1	30-07-2009
			US	2014005361 A1	02-01-2014
			WO	2006000448 A2	05-01-2006

WO 2011064758	A2	03-06-2011	NONE		

WO 2013148117	A1	03-10-2013	CA	2862516 A1	03-10-2013
			WO	2013148117 A1	03-10-2013

WO 2013113008	A1	01-08-2013	AU	2013211846 A1	14-08-2014
			CA	2862745 A1	01-08-2013
			EP	2807266 A1	03-12-2014
			WO	2013113008 A1	01-08-2013
