Title: CATALYST SYSTEM COMPRISING A MIXTURE OF CATALYST COMPONENTS FOR PRODUCING A POLYOLEFIN BLEND

Abstract: Provided is a catalyst system for producing a polyolefin blend, which catalyst system comprises a mixture including the following catalyst components: i) a catalyst component A capable of producing an isotactic olefin polymer, and/or a catalyst component A' capable of producing a polymer comprising an isotactic polyolefin block; and ii) a catalyst component B capable of producing a syndiotactic polyolefin, and/or a catalyst component B' capable of producing a polymer comprising a syndiotactic polyolefin block, wherein each of the components in the catalyst system is distinct from the other components in the catalyst system.
The present invention relates to a catalyst system for use in preparing isotactic polyolefin/syndiotactic polyolefin blends, especially isotactic polypropylene/syndiotactic polypropylene (iPP/sPP) blends. The invention further relates to a catalyst system comprising metallocene catalyst components and a process for producing iPP/sPP blends using the catalyst system.

Olefins having 3 or more carbon atoms can be polymerised to produce a polymer with an isotactic stereochemical configuration. For example, in the polymerisation of propylene to form polypropylene, the isotactic structure is typically described as having methyl groups attached to the tertiary carbon atoms of successive monomeric units on the same side of a hypothetical plane through the main chain of the polymer. This can be described using a three dimensional stereochemical representation and the corresponding Fischer projection formula as follows:

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Me H Me H Me H Me H Me H Me H Me H
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Another way of describing the structure is through the use of NMR spectroscopy. Bovey's NMR nomenclature for an isotactic pentad is "mmmm" with each "m" representing a "meso" diad or successive methyl groups on the same side in the plane.

In contrast to the isotactic structure, syndiotactic polymers are those in which the methyl groups attached to the tertiary carbon atoms of successive monomeric units in the chain lie on alternate sides of the plane of the polymer. The structure of a syndiotactic polymer is described as follows using a three dimensional stereochemical representation and the corresponding Fischer projection formula:
In NMR nomenclature, a syndiotactic pentad is described as "rrrr" in which "r" represents a "racemic" diad with successive methyl groups on alternate sides of the plane.

In contrast to isotactic and syndiotactic polymers, an atactic polymer exhibits no regular order of repeating unit. Unlike syndiotactic or isotactic polymers, an atactic polymer is not crystalline and forms essentially a waxy product.

It is desirable to form blends of isotactic and syndiotactic polyolefins, in particular blends of iPP and sPP. For instance, sPP is generally sticky, which reduces its processibility. This stickiness can be reduced by blending sPP with iPP. In addition, sPP gives high transparency and flexibility in injection blow moulding and injection moulding, whilst iPP gives improved processibility and faster solidification. These advantageous properties can be optimised by blending the two polymers.

Moreover, the softness of polypropylene fibres can be improved by incorporating sPP in iPP, whilst the thermal bonding properties of non-woven fibres can be improved also by incorporating sPP in iPP. By blending a small quantity of sPP with iPP the modulus of polypropylene films can be reduced thus forming films that are less stiff, whilst simultaneously improving the heat shrink, tear resistance and impact resistance properties of the films.

The low crystallisation rate of sPP can be problematic, when processing sPP in a melt. However, the crystallisation rate of iPP is much greater that that of sPP. By
incorporating some iPP in sPP, it is possible to increase the crystallisation rate of sPP in the melt mixture, since as the melt cools, iPP crystallises relatively quickly and serves to seed the crystallisation of sPP. Thus, for this additional reason, it is desirable to form blends of iPP and sPP.

In order to take advantage of the favourable properties of both isotactic and syndiotactic polyolefins, it is known to prepare a physical blend of such polyolefins. However, the properties of the above known physical blends still fall short of the properties which should be achievable from homogeneous reactor blends. This is because polyolefins are generally resistant to blending, in part due to the length of the polymer chains which hinders the intimate intermingling of individual molecules. Furthermore, physical blending is costly, requiring energy to heat the polymer components and to co-extrude the components through a die and sometimes lead to polymer degradation and property changes.

It is an object of the present invention to overcome the above problems associated with the physical blending of isotactic and syndiotactic polyolefins. Accordingly, the present invention provides a catalyst system for producing a polyolefin blend, which catalyst system comprises a mixture including the following catalyst components:

(i) a catalyst component A capable of producing an isotactic olefin polymer, and/or a catalyst component A' capable of producing a polymer comprising an isotactic polyolefin stereoblock; and
(ii) a catalyst component B capable of producing a syndiotactic polyolefin, and/or a catalyst component B' capable of producing a polymer comprising a syndiotactic polyolefin stereoblock;

wherein each of the components in the catalyst system is distinct from the other components in the catalyst system.

In the context of the present invention, catalysts of type A and B mean catalysts capable of producing substantially homopolymeric isotactic polyolefin and substantially homopolymeric syndiotactic polyolefin respectively. Catalysts of type A'
and B' mean catalysts capable of producing block co-polymers of light olefins which comprise predominantly isotactic polyolefin blocks or predominantly syndiotactic polyolefin blocks respectively. The further blocks in the block co-polymers are not especially limited and may be blocks formed from olefin monomers, or blocks formed from other monomers.

The catalyst system of the present invention comprises at least two different catalyst components, one component of type A (and/or A') and one component of type B (and/or B'). Thus, the present catalyst system may comprise any of the following component combinations:

A+B  
A+B'  
A'+B  
A'+B'  
A+B+A'  
A+B+B'  
A+B'+A'  
A+B+B'  ; and  
A+A'+B+B'  

as well as further catalyst components, if desired.

In the case where a catalyst component is capable of producing an olefin polymer comprising both isotactic and syndiotactic blocks, then this catalyst component is simultaneously of component type A' and component type B' and in the present context is termed an A'B' component. However, for the avoidance of doubt, the present system cannot comprise one A'B' component alone, since the component of type (i) should be a different catalyst from the component of type (ii) in order that the catalyst system is capable of producing at least two substantially different polyolefins. However, the present catalyst system may comprise two different A'B' components.
The present invention further provides a method for producing a blend of an isotactic polyolefin and a syndiotactic polyolefin, which method comprises polymerising an olefin monomer in the presence of a catalyst system as defined above.

The use of a catalyst mixture to produce an in situ mixture of isotactic polyolefin and syndiotactic polyolefin (a chemical blending process) leads to polymer blends (chemical blends) that are much more homogeneous than known blends produced by a physical blending process. The improved homogeneity of the present blends contributes to their improved physical and mechanical properties.

Thus, the polymer blends produced by the catalyst system of the present invention have improved processibility and solidification properties in injection blow moulding and injection moulding, whilst simultaneously showing high transparency and flexibility. Fibres formed from the present blends have improved softness and improved thermal bonding characteristics. Films produced from the present blends have a reduced modulus, being less stiff and have improved heat shrink, tear resistance and impact resistance characteristics.

Typically, in the present catalyst system catalysts components (i) and (ii) are metallocene compounds. However, the catalyst components (i) and (ii) used in the present catalyst system for producing the iPP/sPP blend are not particularly limited, provided that they can be mixed with each other.

In a preferred embodiment of the present invention, the catalyst component (i) comprises a catalyst A having the following formula:

\[ R''(CpR^1R^2)(Cp'R^'_n)MQ_2 \]

wherein Cp is a substituted cyclopentadienyl ring; Cp' is a substituted or unsubstituted fluorenyl ring; R'' is a structural bridge imparting stereorigidity to the component; R^1 is a substituent on the cyclopentadienyl ring which is distal to the bridge, which distal substituent comprises a bulky group of the formula XR*3 in
which X is an atom from group IVA and each R* is the same or different and is
chosen from a hydrogen or a hydrocarbyl group having from 1-20 carbon atoms, R^2
is a substituent on the cyclopentadienyl ring which is proximal to the bridge and
positioned non-vicinal to the distal substituent and is a hydrogen or is of the formula
YR#_3 in which Y is an atom from group IVA, and each R# is the same or different
and is chosen from a hydrogen or a hydrocarbyl group having from 1-7 carbon
atoms, each R' is the same or different and is a hydrocarbyl group having from 1-20
carbon atoms, and n is an integer of from 0-8; M is a metal atom from group IVB or is
vanadium; and each Q is a hydrocarbon having from 1-20 carbon atoms or is a
halogen;

and/or a catalyst A having the following formulae:

(Ind)_2R'MQ_2 or;

(IndH_4)_2R''MQ_2

wherein each Ind is the same or different and is a substituted or unsubstituted
indenyl group, wherein each IndH_4 is the same or different and is a substituted or
unsubstituted tetrahydroindenyl group, R' is a structural bridge imparting
stereorigidity to the component, R'' is a structural bridge imparting stereorigidity to
the component; M is a metal atom from group IVB or is vanadium; and each Q is a
hydrocarbon having from 1-20 carbon atoms or is a halogen. Preferably the
structural bridge comprises a C_1-C_4 alkylene group.

In common with catalyst A, the catalyst B used in component (ii) of the present
catalyst system for producing the syndiotactic polyolefin is not particularly limited,
provided that it can be mixed with catalyst component (i). It is preferred in the
present invention that the catalyst B is a catalyst having the following formula:

R''(CpR_m)(Cp'R'_n)MQ_2

wherein Cp is a substituted or unsubstituted cyclopentadienyl ring; Cp' is a
substituted or unsubstituted fluorenyl ring; R'' is a structural bridge imparting
stereorigiditiy to the component; each R is the same or different and is a hydrocarbyl group having from 1-20 carbon atoms; each R’ is the same or different and is a hydrocarbyl group having from 1-20 carbon atoms; m is an integer of from 0-4; r is an integer from 0-8; M is a metal atom from group IVB or is vanadium; and each Q is a hydrocarbon having from 1-20 carbon atoms or is a halogen. Preferably the CpR_m group possesses bilateral symmetry and more preferably is unsubstituted (m = 0).

Preferred catalysts of type A’ are those having the following formula:

\[ R''(CpR_x)(Cp'R'_y)MO_2 \]

wherein Cp is a substituted cyclopentadienyl ring; Cp’ is a substituted or unsubstituted fluorenyl ring; R” is a structural bridge imparting stereoriditiy to the component; each R is the same or different and is a hydrocarbyl group having from 1-20 carbon atoms, each R’ is the same or different and is a hydrocarbyl group having from 1-20 carbon atoms, and x and y are independently an integer of from 0-4 and 0-8 respectively; M is a metal atom from group IVB or is vanadium; and each Q is a hydrocarbon having from 1-20 carbon atoms or is a halogen; wherein the CpR_x group lacks bilateral symmetry. In a preferred embodiment, the Cp group is substituted at the 3-position. Particularly preferably the substituent comprises a trimethylsilyl group.

Preferred catalysts of type B’ are those having the following formula:

\[ R''(CpR_q)XMQ \]

wherein Cp is a substituted or unsubstituted cyclopentadienyl ring or a substituted or unsubstituted fluorenyl ring; R” is a structural bridge between Cp and X imparting stereoriditiy to the component; each R is the same or different and is selected from a hydrocarbyl group having from 1-20 carbon atoms, a halogen, an alkoxy group, an alkoxyalkyl group, an alkylamino group or an alkylsilylo group; when Cp is a cyclopentadienyl ring; q is an integer from 0-4; and when Cp is a fluorenyl ring q is
an integer from 0-8; X is a heteroatom from group VA or group VIA; M is a metal atom from group IIIb, IVB, VB or VIB in any of its theoretical oxidation states; and each Q is a hydrocarbon having from 1-20 carbon atoms or is a halogen; wherein the bilateral symmetry of the CpR_Q group is maintained. Thus, preferably the CpR_Q group is symmetrically substituted.

The substituent or substituents present on the cyclopentadiene, indene and fluorine rings in the above-described catalysts are not particularly limited. The above rings, when comprising more than one substituent, may be substituted with the same substituent throughout, or with different substituents. Typically the substituents are independently selected from an aryl group and a hydrocarbyl group having from 1-20 carbon atoms. The most preferred substituents are methyl groups. Other preferred substituents include Et, n-Pr, i-Pr, n-Bu, t-Bu, Me_3Si, R-O, cycloalkyl, and halogen.

The type of bridge present between the rings in the above-described catalysts is not itself particularly limited. Typically R'' comprises an alkylidene group having 1 to 20 carbon atoms, a germanium group (e.g. a dialkyl germanium group), a silicon group (e.g. a dialkyl silicon group), a siloxane group (e.g. a dialkyl siloxane group), an alkyl phosphine group or an amine group. Preferably, the substituent comprises a silyl radical or a hydrocarbyl radical having at least one carbon atom to form the bridge, such as a substituted or unsubstituted ethylenyl radical (e.g. -CH_2CH_2-). Most preferably R'' is isopropyldene (Me_2C), Ph_2C, ethylenyl, or Me_2Si. It is particularly preferred that catalyst components comprising a bisindenyl moiety comprise an ethylenyl or an Me_2Si bridge, whilst catalyst components comprising a cyclopentadienyl-fluorenyl moiety comprise an Me_2C, Ph_2C, or Me_2Si bridge.

Some specific examples of preferred catalysts according to the present invention are the following:

A (iPP catalysts):

Me_2Si(2-Me-Benz-Ind)_2 ZrCl_2
Me₂Si(2-Me-4-Naphthyl-Ind)₂ ZrCl₂
Me₂Si(2-Me-Ind)₂ ZrCl₂
Ph₂C(t-BuCp)(Flu) ZrCl₂
Et(Ind)₂ ZrCl₂
Et(IndH₄)₂ ZrCl₂
Me₂Si(2-Me-4,5Benzyl-Ind)₂ ZrCl₂
(R1R2Cp-Flu)RZrCl₂
A' (iPP block catalysts)
Ph₂C(3-Me₃Si-Cp)(Flu) ZrCl₂

B (sPP catalysts)
Ph₂C(Cp)(Flu) ZrCl₂

B' (sPP block catalysts)
Me₂S(t-BuN)(2,7-di-t-Bu-Flu) ZrCl₂

The catalyst components (i) and (ii) of the present catalyst system can be mixed by a physical process (physical blending), such as by slurrying a powdered form of the catalysts together in a hydrocarbon solvent. Alternatively components (i) and (ii) can be mixed by a chemical process (chemical blending). Chemical blending can be effected by, for example, forming a solution containing both catalysts (i) and (ii) and adding the solution to the support. It is preferred that chemical blending is carried out by immobilising both catalyst (i) and catalyst (ii) on a solid support in such a manner that one or more individual particles of the solid support have both catalyst (i) and catalyst (ii) immobilised on them.

The proportion of catalyst (i) to catalyst (ii) in the present catalyst system is not especially limited, and depends on the ratio of isotactic to syndiotactic polymer required in the final product. This will be dependent on the particular application for which the product is to be used. Typically, the proportion of catalyst (i) to catalyst (ii) is such that the final isotactic/syndiotactic blend comprises 50 wt. % or more isotactic
polyolefin and from 0.5-50 wt. % of syndiotactic polyolefin. More preferably, the final blend comprises from 0.3-15 wt. % of syndiotactic polyolefin and most preferably from 1-10 wt. % of syndiotactic polyolefin.

In order to produce polyolefin blends displaying an especially high degree of homogeneity (and consequently especially favourable properties) it is preferred that the present catalyst system comprises a catalyst component capable of producing a polyolefin comprising both isotactic polyolefin blocks and syndiotactic or atactic polyolefin blocks (an A' catalyst), or a catalyst capable of producing a polyolefin comprising both syndiotactic polyolefin blocks and isotactic or atactic polyolefin blocks (a B' catalyst). In the context of the present invention, these catalysts are termed stereoblock catalysts. Preferred catalyst components of this type are those capable of producing a polyolefin comprising both isotactic and syndiotactic polyolefin blocks (i.e. A'B' catalyst components). It is particularly preferred that the present catalyst system comprises an A component, a B component and an A'B' component.

Without being bound by theory, it is believed that, since the polyolefins produced by the stereoblock catalysts comprise both isotactic and syndiotactic units, they mix more readily with isotactic and syndiotactic homopolymers than these homopolymers do with each other. Thus, the inclusion of one or more stereoblock catalysts in the present catalyst system improves miscibility, i.e. it ensures that the polyolefin blend produced comprises a polymer that promotes mixing between the isotactic and syndiotactic polyolefins. The resulting polymer blend is thus more homogeneous than would otherwise be the case. Consequently the polymer has fewer weak spots in the crystal structure, leading to significant improvements in its mechanical strength.

In addition to the above catalyst components (i) and (ii), the catalyst system of the present invention may comprise one or more co-catalysts capable of activating any one or more of the catalyst components. Typically, the co-catalyst comprises an aluminium- or boron-containing co-catalyst.
Suitable aluminium-containing co-catalysts comprise an alumoxane, an alkyl aluminium compound and/or a Lewis acid.

The alumoxanes that can be used in the present invention are well known and preferably comprise oligomeric linear and/or cyclic alkyl alumoxanes represented by the formula (I):

$$R\left\{\frac{\text{Al-O}}{n}\right\}_{n} R_2$$

for oligomeric linear alumoxanes; and formula (II)

$$\left\{\frac{\text{Al-O}}{m}\right\}_m$$

(II)

for oligomeric cyclic alumoxanes,

wherein $n$ is 1-40, preferably 10-20; $m$ is 3-40, preferably 3-20; and R is a C$_1$-C$_8$ alkyl group, preferably methyl. Generally, in the preparation of alumoxanes from, for example, aluminium trimethyl and water, a mixture of linear and cyclic compounds is obtained.

Suitable boron-containing co-catalysts may comprise a triphenylcarbenium boronate, such as tetrakis-pentafluorophenyl-borato-triphenylcarbenium as described in EP-A-0427696:
The catalyst system of the present invention may be employed in any method, provided that the required catalytic activity is not impaired. In a preferred embodiment of the present invention, the catalyst system is employed in a solution polymerisation process, which is homogeneous, or a slurry process, which is heterogeneous. In a solution process, typical solvents include hydrocarbons having 4-7 carbon atoms such as heptane, toluene or cyclohexane. In a slurry process it is necessary to immobilise the catalyst system on an inert support, particularly a porous solid support such as talc, inorganic oxides and resinous support materials such as polyolefin. Preferably, the support material is an inorganic oxide in its finely divided form.

Suitable inorganic oxide materials which are desirably employed in accordance with this invention include group IIA, IIIA, IVA, or IVB metal oxides such as silica, alumina and mixtures thereof. Other inorganic oxides that may be employed either alone or in combination with the silica, or alumina are magnesia, titania, zirconia, and the like. Other suitable support materials, however, can be employed, for example, finely divided functionalised polyolefins such as finely divided polyethylene.
Preferably, the support is a silica support having a surface area of from 200-700 m²/g and a pore volume of from 0.5-3 ml/g.

The amount of alumoxane and metalloccenes usefully employed in the preparation of the solid support catalyst can vary over a wide range. Preferably the aluminium to transition metal mole ratio is in the range 1:1 to 130:1, preferably 1:1 to 100:1, more preferably 5:1 to 70:1 and most preferably 5:1 to 50:1.

The order of addition of the catalyst components and alumoxane to the support material can vary. In accordance with a preferred embodiment of the present invention alumoxane dissolved in a suitable inert hydrocarbon solvent is added to the support material slurried in the same or other suitable hydrocarbon liquid and thereafter a mixture of the catalyst components is added to the slurry.

Preferred solvents include mineral oils and the various hydrocarbons which are liquid at reaction temperature and which do not react with the individual ingredients. Illustrative examples of the useful solvents include the alkanes such as pentane, isopentane, hexane, heptane, octane and nonane; cycloalkanes such as cyclopentane and cyclohexane, and aromatics such as benzene, toluene, ethylbenzene and diethylbenzene.

Preferably the support material is slurried in toluene and the catalyst components and alumoxane are dissolved in toluene prior to addition to the support material.

The polyolefins that catalysts (i) and (ii) are capable of producing are not particularly limited, except that the monomeric olefin unit forming each polyolefin must have three or more carbon atoms. It is particularly preferred that both of catalyst components (i) and (ii) are capable of producing polypropylene.

The catalyst system of the present invention is used in the method of the present invention to produce blends of isotactic polyolefin and syndiotactic polyolefin. It is especially preferred that the method of the present invention is a method of
producing a blend of isotactic polypropylene and syndiotactic polypropylene (an iPP/sPP blend).

The conditions employed for polymerisation in the method of the present invention are not particularly limited, provided they are sufficient to effectively polymerise the particular monomeric olefin used as a starting material. Typical polymerisation conditions in a slurry polymerisation are at a temperature of from 20-120°C, a pressure of from 0.1-5.6 MPa and a reaction time of from 10 mins to 4 hours.

The polyolefin blends of the present invention, and in particular the iPP/sPP, blends may be used to produce fibres. For the production of spunlaid iPP/sPP fibres, a typical extrusion temperature is in the range of from 200-260°C, most typically from 230-250°C. For the production of staple fibres, a typical extrusion temperature would be in the range of from 230-330°C, most typically from 280-300°C.

Fibres produced in accordance with the present invention may be produced from iPP/sPP blends having other additives to improve the mechanical processing or spinnability of the fibres. The fibres may be used to produce non-woven fabrics for use in filtration; in personal care products such as wipers, diapers, feminine hygiene products and incontinence products; in medical products such as wound dressings, surgical gowns, bandages and surgical drapes; in protective covers; in outdoor fabrics and in geotextiles. Non-woven fabrics made with iPP/sPP fibres can be part of such products, or constitute entirely the products. As well as making non-woven fabrics, the fibres may also be employed to make a woven knitted fabric or mat. The non-woven fabrics produced from the fibres in accordance with the invention can be produced by several processes, such as air through blowing, melt blowing, spun bonding or bonded carded processes. The fibres may also be formed as a non-woven spunlace product which is formed without thermal bonding by fibres being entangled together to form a fabric by the application of a high pressure fluid such as air or water.
The invention will now be described in further detail by way of example only, with reference to the following specific embodiments.

Examples

Example 1
Catalyst preparation
(Me₂Si)(2-Me-4,5-Benzyl-Ind)₂ZrCl₂ (a catalyst A) and (Ph₂C)(Cp)(Flu)ZrCl₂ (a catalyst B) were prepared.

Catalyst A preparation – two steps procedure
The support used was silica having a total pore volume of 4.22 ml/g and a surface area of 322 m²/g. This silica was dried in a fluidised bed reactor (for 6 hours at 150°C with 75 Nl of H₂) to remove the physically absorbed water.

5 g of this dried silica were suspended in 100 ml of dried toluene in a round bottom flask equipped with a magnetic stirrer, a nitrogen inlet and a dropping funnel. A 30wt% MAO solution in toluene (0.7-1.3 weight equivalents of MAO to silica, preferably 0.9 weight equivalents) was added dropwise to the silica suspension at room temperature. A reaction between MAO and the hydroxyl groups of the silica occurred, and the exothermic reaction (10 °C temperature rise) was accompanied by methane gas release. The slurry was heated to 110°C and allowed to react for 4 hours at toluene reflux. The suspension was then filtered on a frit funnel. The reaction product was washed until the filtrate was substantially free of MAO. After the washing with toluene, the precipitate was then washed with pentane to facilitate
drying. The support was finally dried under a mild vacuum. 9.7 g of support was obtained as a white powder.

To produce the activated catalyst, the metallocene (Me₂Si)(2-Me-4,5-Benzyl-Ind)₂ZrCl₂ (2 to 6 wt% of metallocene loading on the final catalyst, preferably 4 wt%) was dissolved in toluene and added to the toluene suspended reaction product of silica and MAO at room temperature. Reaction was allowed to take place for 2 hours at room temperature to form the active sites by reaction of the remaining aluminium alkyl functions on the support with the metallocene. The suspension was filtered and washed with toluene until filtrate was colourless. The precipitate was then washed with pentane to facilitate drying and finally dried under mild vacuum.

* Catalyst B preparation – one pot procedure.*

The support used was silica having a total pore volume of 4.22 ml/g and a surface area of 322 m²/g. This silica was dried in a fluidised bed reactor (for 6 hours at 150°C with 75 Nl of H₂) to remove the physically absorbed water.

5 g of this dried silica were suspended in 100 ml of dried toluene in a round bottom flask equipped with a magnetic stirrer, a nitrogen inlet and a dropping funnel. In this procedure, the ion pair was formed by initial reaction of MAO solution with the metallocene (Ph₂C)(Cp)(Flu)ZrCl₂ (2 to 6 wt% of metallocene to final catalyst, preferably 6 wt%). The ion pair solution was then added to the silica/toluene suspension (0.7-1.3 weight equivalents of MAO to silica, preferably 0.9 wt equivalents). The suspension was allowed to react for 4 hours at toluene reflux. The resulting slurry was filtered and washed with toluene and pentane and finally dried under mild vacuum.

*Physical blends of catalysts preparation*

To produce a physical blend of supported catalysts, powder of each supported metallocene catalyst was mixed to produced 75:25, 83:17 and 90:10 physical blends.
of catalyst A:catalyst B. In each case, the catalyst system comprised a 6 wt.% loading of metallocene.

**Example 2**

*Catalyst preparation*

(Me$_2$Si)(2-Me-Ind)$_2$ZrCl$_2$ (a catalyst A) and (Ph$_2$C)(Cp)(Flu)ZrCl$_2$ (a catalyst B) were prepared.

*Catalyst A preparation – two steps procedure*

The catalyst system was prepared according to the two steps procedure described above in Example 1.

*Catalyst B preparation – one pot procedure*

The catalyst system was prepared according to the one pot procedure described above in Example 1.

*Physical blends of catalysts preparation*

A physical blend of supported catalysts was produced by mixing powder of each supported metallocene catalyst to produce a 72:25 physical blend of catalyst A:catalyst B. In each case, the catalyst system comprised a 6 wt.% loading of metallocene.

**Example 3**

*Catalyst preparation – two steps procedure*

(Ph$_2$C)((Me$_3$Si)Cp)(Flu)ZrCl$_2$ (a catalyst A') and (Ph$_2$C)(Cp)(Flu)ZrCl$_2$ (a catalyst B) were prepared.

The support used was silica having a total pore volume of 4.22 ml/g and a surface area of 322 m$^2$/g. This silica was dried in a fluidised bed reactor (for 6 hours at 150°C with 75 Nl of H$_2$) to remove the physically absorbed water.
5 g of this dried silica were suspended in 100 ml of dried toluene in a round bottom flask equipped with a magnetic stirrer, a nitrogen inlet and a dropping funnel. A 30 wt.% MAO solution in toluene (0.7-1.3 weight equivalents of MAO to silica, preferably 0.9 weight equivalents) was added dropwise to the silica suspension at room temperature. A reaction between MAO and the hydroxyl groups of the silica occurred, and the exothermic reaction (10°C temperature rise) was accompanied by methane gas release. The slurry was heated to 110°C and allowed to react for 4 hours at toluene reflux. The suspension was then filtered on a frit funnel. The reaction product was washed until the filtrate was substantially free of MAO. After the washing with toluene, the precipitate was then washed with pentane to facilitate drying. The support was finally dried under a mild vacuum. 9.7 g of support was obtained as a white powder.

To produce the activated catalyst, the metallocenes (Ph₂C)((Me₃Si)Cp)(Flu)ZrCl₂ and (Ph₂C)(Cp)(Flu)ZrCl₂ were dissolved in toluene and added to toluene suspended reaction product of silica and MAO at room temperature. Reaction was allowed to take place for 2 hours at room temperature to form the active sites by reaction of the remaining aluminium alkyl functions on the support with the metallocene. The suspension was filtered and washed with toluene until filtrate was colourless. The precipitate was then washed with pentane to facilitate drying and finally dried under mild vacuum to produce an 80:20 (by weight) chemical blend of catalyst A:catalyst B. This procedure was repeated to produce 90:10 and 95:5 chemical blends of A:B. The total loading of metallocenes was 6 wt.%.

Polymerisation of propylene
Each of the above catalyst systems, as well as systems comprising only the individual catalyst components, were used to polymerise propylene. 2 litres of liquid propylene were polymerised using 100 mg of each catalyst.
Example 1 - Table 1 below shows the results of microtacticity analysis from $^{13}$C NMR data in which the percent of mmmm pentads decreases as the content of Benzyl-Ind catalyst in the catalyst system decreases.

<table>
<thead>
<tr>
<th>Catalyst</th>
<th>Wt.% sPP</th>
<th>% mmmm</th>
<th>% rrrr</th>
<th>% m</th>
<th>% r</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>iPP and sPP catalysts alone</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Benzyl-Ind (iPP)</td>
<td>---</td>
<td>93.7</td>
<td>0.37</td>
<td>97.4</td>
<td>2.6</td>
</tr>
<tr>
<td>Cp (sPP)</td>
<td>---</td>
<td>2.4</td>
<td>75.4</td>
<td>10.2</td>
<td>89.8</td>
</tr>
<tr>
<td><strong>Physical blends of iPP and sPP catalysts</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Benzyl-Ind/Cp; 75/25</td>
<td>28.3</td>
<td>69.3</td>
<td>19.2</td>
<td>74.9</td>
<td>25.1</td>
</tr>
<tr>
<td>Benzyl-Ind/Cp; 83/17</td>
<td>21.7</td>
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<td>13.2</td>
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<td>87.6</td>
<td>3.7</td>
<td>92.5</td>
<td>7.5</td>
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Example 2 - Table 2 below shows the results of microtacticity analysis from $^{13}$C NMR data in which the percent of mmmm pentads decreases as the content of Me-Ind catalyst in the catalyst system decreases.

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<tr>
<th>Catalyst</th>
<th>Wt.% sPP</th>
<th>% mmmm</th>
<th>% rrrr</th>
<th>% m</th>
<th>% r</th>
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<tr>
<td><strong>iPP and sPP catalysts alone</strong></td>
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<td></td>
<td></td>
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<tr>
<td>Me-Ind (iPP)</td>
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<td>93.4</td>
<td>0.0</td>
<td>97.4</td>
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<tr>
<td>Cp (sPP)</td>
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<td>2.4</td>
<td>75.4</td>
<td>10.2</td>
<td>89.8</td>
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<tr>
<td><strong>Physical blend of iPP and sPP catalysts</strong></td>
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<tr>
<td>Me-Ind/Cp; 75/25</td>
<td>---</td>
<td>78.4</td>
<td>12.3</td>
<td>83.4</td>
<td>16.6</td>
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Example 3 - Table 3 below shows the results of microtacticity analysis from $^{13}$C NMR data in which, for both the chemically and physically blended catalyst systems,
the percent of mmmm pentads increases as the content of TMSCp catalyst in the catalyst system increases.

Table 3

<table>
<thead>
<tr>
<th>Catalyst</th>
<th>Wt.% iPP in sPP</th>
<th>% mmmm</th>
<th>% rrrr</th>
<th>% m</th>
<th>% r</th>
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<tbody>
<tr>
<td><strong>iPP/sPP and sPP catalysts alone</strong></td>
<td></td>
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<td></td>
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<tr>
<td>TMSCp (IPP/sPP)</td>
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<td>53.8</td>
<td>17.5</td>
<td>69.9</td>
<td>30.1</td>
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<tr>
<td>Cp (sPP)</td>
<td>---</td>
<td>2.4</td>
<td>75.4</td>
<td>10.2</td>
<td>89.8</td>
</tr>
<tr>
<td><strong>Physical blends of iPP/sPP and sPP catalysts</strong></td>
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<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>TMSCp/Cp; 50/50</td>
<td>1.0</td>
<td>3.0</td>
<td>75.0</td>
<td>10.8</td>
<td>89.2</td>
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<tr>
<td>TMSCp/Cp; 75/25</td>
<td>8.5</td>
<td>6.8</td>
<td>70.5</td>
<td>15.2</td>
<td>84.8</td>
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<tr>
<td>TMSCp/Cp; 90/10</td>
<td>22.3</td>
<td>13.9</td>
<td>63.1</td>
<td>23.5</td>
<td>76.5</td>
</tr>
<tr>
<td><strong>Chemical blends of iPP/sPP and sPP catalysts</strong></td>
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<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>TMSCp/Cp; 80/20</td>
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<td>22.2</td>
<td>77.8</td>
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<tr>
<td>TMSCp/Cp; 95/5</td>
<td>30.0</td>
<td>17.5</td>
<td>57.7</td>
<td>28.1</td>
<td>71.9</td>
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CLAIMS:

1. A catalyst system for producing a polyolefin blend, which catalyst system comprises a mixture including the following catalyst components:

(i) a catalyst component A capable of producing an isotactic olefin polymer, and/or a catalyst component A' capable of producing a polymer comprising an isotactic polyolefin block; and
(ii) a catalyst component B capable of producing a syndiotactic polyolefin, and/or a catalyst component B' capable of producing a polymer comprising a syndiotactic polyolefin block;

wherein each of the components in the catalyst system is distinct from the other components in the catalyst system, and

wherein the catalyst component A has the formula:

\[ R''(CpR^1R^2)(Cp'R_n')MQ_2 \]

wherein Cp is a substituted cyclopentadienyl ring; Cp' is a substituted or unsubstituted fluorenyl ring; R'' is a structural bridge imparting stereorigidity to the component; R^1 is a substituent on the cyclopentadienyl ring which is distal to the bridge, which distal substituent comprises a bulky group of the formula XR^*_3 in which X is an atom from group IVA and each R^* is the same or different and is chosen from a hydrogen or a hydrocarbaryl group having from 1-20 carbon atoms, R^2 is a substituent on the cyclopentadienyl ring which is proximal to the bridge and positioned non-vicinal to the distal substituent and is a hydrogen or is of the formula YR^#_3 in which Y is an atom from group IVA, and each R^# is the same or different and is chosen from a hydrogen or a hydrocarbaryl group having from 1-7 carbon atoms, each R' is the same or different and is a hydrocarbaryl group having from 1-20 carbon atoms, and n is an integer of from 0-8; M is a metal atom from group IVB or is vanadium; and each Q is a hydrocarbon having from 1-20 carbon atoms or is a halogen;
and/or has the formula:

$$(\text{Ind})2R'MQ2 \text{ or;}$$

$$(\text{IndH}_4)2R''MQ2$$

wherein each Ind is the same or different and is a substituted or unsubstituted indenyl group, wherein each IndH$_4$ is the same or different and is a substituted or unsubstituted tetrahydroindenyl group, R' is a structural bridge imparting stereorigidity to the component, R'' is a structural bridge imparting stereorigidity to the component; M is a metal atom from group IVB or is vanadium; and each Q is a hydrocarbon having from 1-20 carbon atoms or is a halogen, and wherein the catalyst component B has the formula:

$$R''(\text{CpR}_M)(\text{Cp'R}_r)MQ_2$$

wherein Cp is a substituted or unsubstituted cyclopentadienyl ring; Cp' is a substituted or unsubstituted fluorenyl ring; R'' is a structural bridge imparting stereorigidity to the component; each R is the same or different and is a hydrocarbyl group having from 1-20 carbon atoms; each R' is the same or different and is a hydrocarbyl group having from 1-20 carbon atoms; m is an integer of from 0-4; r is an integer from 0-8; M is a metal atom from group IVB or is vanadium; and each Q is a hydrocarbon having from 1-20 carbon atoms or is a halogen, and wherein the catalyst component A' has the formula:

$$R''(\text{CpR}_X)(\text{Cp'R}_Y)MQ_2$$

wherein Cp is a substituted cyclopentadienyl ring; Cp' is a substituted or unsubstituted fluorenyl ring; R'' is a structural bridge imparting stereorigidity to the component; each R is the same or different and is a hydrocarbyl group having from 1-20 carbon atoms, each R' is the same or different and is a hydrocarbyl group having from 1-20 carbon atoms, and x and y are independently an integer of from 0-4 and 0-8 respectively; M is a metal atom from group IVB or is vanadium; and each Q is a hydrocarbon having from 1-20 carbon atoms or is a halogen; wherein the
CpR_X group lacks bilateral symmetry; and wherein the Cp group is preferably substituted at the 3-position, and wherein the catalyst component B' has the formula:

\[ R''(\text{CpR}_q)\text{XMQ} \]

wherein Cp is a substituted cyclopentadienyl ring or a substituted or unsubstituted fluorenyl ring; R'' is a structural bridge between Cp and X imparting stereorigidity to the component; each R is the same or different and is selected from a hydrocarbyl group having from 1-20 carbon atoms, a halogen, an alkoxy group, an alkoxyalkyl group, an alkylamino group or an alkylsilylo group; when Cp is a cyclopentadienyl ring; q is an integer from 0-4; and when Cp is a fluorenyl ring q is an integer from 0-8; X is a heteroatom from group VA or group VIA; M is a metal atom from group IIIB, IVB, VB or VIB in any of its theoretical oxidation states; and each Q is a hydrocarbon having from 1-20 carbon atoms or is a halogen; wherein the bilateral symmetry of the CpR_q group is maintained; and wherein the CpR_q group is preferably symmetrically substituted.

2. A catalyst system according to claim 1, wherein M is Ti, Zr, or Hf.

3. A catalyst system according to claim 1 or claim 2, wherein Q is Cl.

4. A catalyst system according to any one of the preceding claims, wherein R'' is substituted or unsubstituted and comprises an alkylene radical having from 1-20 carbon atoms and preferably from 1-4 carbon atoms, a dialkyl germanium group, a dialkyl silicon group, a dialkyl siloxane group, an alkyl phosphine radical or an amine radical.

5. A catalyst system according to any one of the preceding claims, wherein R'' comprises an isopropylidene, (Me_2C), Ph_2C, ethylenyl, Me_2S or Me_2Si group.
6. A catalyst system according to anyone of the preceding claims, comprising a physical mixture of catalyst components.

7. A catalyst system according to any one of the preceding claims, comprising a chemical mixture of catalyst components.

8. A catalyst system according to any one of the preceding claims, wherein catalyst component A and/or A' and/or catalyst component B and/or B' are immobilised on a solid support.

9. A catalyst system according to any one of the preceding claims, further comprising an aluminium- or boron-containing activating agent capable of activating catalyst component A and/or A' and/or catalyst component B and/or B'.

10. A catalyst system according to any one of the preceding claims, wherein catalyst component A is capable of producing isotactic polypropylene and/or catalyst component B is capable of producing syndiotactic polypropylene.

11. A method for producing a blend of an isotactic polyolefin and a syndiotactic polyolefin, which method comprises polymerising an olefin monomer in the presence of a catalyst system as defined in any of claims 1-10.

12. A method according to claim 11, wherein the olefin monomer is propylene.

17. A polyolefin product comprising a blend of an isotactic polyolefin and a syndiotactic polyolefin, obtainable according to a method as defined in claim 11 or claim 12.

18. Use of a catalyst system as defined in any one of claims 1 to 10 for the preparation of a substantially homogenous isotactic polyolefin/syndiotactic polyolefin blend.
## INTERNATIONAL SEARCH REPORT

### A. CLASSIFICATION OF SUBJECT MATTER

**IPCl** 7 C08F4/64 C08F10/06 C08F10/00

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)

**IPCl** 7 C08F

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 practical, search terms used)

EPO-Internal

### C. DOCUMENTS CONSIDERED TO BE RELEVANT

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<th>Citation of document, with indication, where appropriate, of the relevant passages</th>
<th>Relevant to claim No.</th>
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<tr>
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<td>EP 0 870 779 A (FINA TECHNOLOGY) 14 October 1998 (1998-10-14) claims 1,8-11,16</td>
<td>1-5, 7-12,17, 18</td>
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<td>US 5 714 427 A (DOLLE VOLKER ET AL) 3 February 1998 (1998-02-03)</td>
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<td>examples 2,3, column 7, line 55 - line 58</td>
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Further documents are listed in the continuation of box C.

Patent family members are listed in annex.

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Date of the actual completion of the international search 23 January 2003

Date of mailing of the international search report 05/02/2003

Name and mailing address of the ISA

European Patent Office, P.O. 5618 Patentlaan 2 NL - 2280 HV Rijswijk
Tel. (+31-70) 340-2040, Tx 31 651 epo nl, Fax (+31-70) 340-3016

Authorized officer

Fischer, B

Form PCT/ISA/210 (second sheet) (July 1992)
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