(54) Title: FUNCTIONALIZED PARTICULATE BICARBONATE AS BLOWING AGENT, FOAMABLE POLYMER COMPOSITION CONTAINING IT, AND ITS USE IN MANUFACTURING A THERMOPLASTIC FOAMED POLYMER

(57) Abstract: A chemical blowing agent for foaming a thermoplastic polymer, for example PVC plastisol or a polymer resin in an extrusion process, said chemical blowing agent comprising a functionalized particulate bicarbonate containing at least one polyoxyalkylene additive, preferably excluding an exothermic blowing agent. The polyoxyalkylene additive may be selected from the group consisting of polyoxyalkylenes, any derivative thereof, salts thereof, and any combinations thereof. The particulate bicarbonate may be preferably functionalized by spray-coating, extrusion or co-grinding with at least one polyoxyalkylene additive. The functionalized particulate bicarbonate may comprise 50 wt% to less than 100 wt% of the bicarbonate component, and 0.02-50 wt% of the additive. The functionalized particulate bicarbonate may further comprise 0.1 - 5 wt% silica. A foamy polymer composition comprising such chemical blowing agent. A process for manufacturing a foamed polymer, such as foamed PVC, comprising shaping and heating the foamy polymer composition, and a foamed polymer obtained by such process.
FUNCTIONALIZED PARTICULATE BICARBONATE AS BLOWING AGENT, FOAMABLE POLYMER COMPOSITION CONTAINING IT, AND ITS USE IN MANUFACTURING A THERMOPLASTIC FOAMED POLYMER

CROSS-REFERENCE TO RELATED APPLICATIONS
The present application claims priority to European application No. 1718241.3 filed July 20, 2017 and European application No. 18152755.7 filed January 22, 2018, the whole content of each of these applications being incorporated herein by reference for all purposes.

STATEMENT REGARDING FEDERALLY SPONSORED RESEARCH OR DEVELOPMENT
Not applicable.

TECHNICAL FIELD OF THE INVENTION
The invention relates to a functionalized particulate bicarbonate containing an additive. The invention also relates to a foamable composition containing it, such as a PVC plastisol, and to its use/method for manufacturing a thermoplastic foamed polymer, in particular foamed PVC.

BACKGROUND
Polymer foams are found virtually everywhere in our modern world and are used in a wide variety of applications such as disposable packaging of fast-food, the cushioning of furniture and insulation material.

Polymer foams are made up of a solid and gas phase mixed together to form a foam. By combining the two phases fast results in foaming and forming a polymer matrix with either gas bubbles or gas tunnels incorporated in it, which is known as either closed-cell or open-cell structure. Closed-cell foams are generally more rigid, while open-cell foams are usually flexible.

The gas that is used in the foam is termed a blowing agent, and can be either chemical or physical. Chemical blowing agents are chemicals that take part in a reaction or decompose, giving off a gas in the process. Physical blowing agents are gases that do not react chemically in the foaming process and are therefore inert to the polymer forming the matrix.
For the processing of thermoplastic materials, such as polyvinyl chloride (PVC) or polyolefins (PO, PE, PP), styrenics (PS, ABS, ASA, SAN) and natural and synthetic rubber such as nitrile butadiene rubber (NBR) or chloroprene rubber (CR), chemical blowing agents are used for decades. Chemical blowing agents are additives in the manufacturing of foamed thermoplastic polymers. Chemical blowing agents are stable at room temperature but decompose at elevated temperatures during the processing of the polymers while generating gas. This gas creates a foam structure in the thermoplastic polymer. Chemical blowing agents are used in a wide variety of applications including the production of foamed wall papers, artificial leather, floor and wall coverings, carpet backings, thermal insulation materials, insulation sealants, footwear, automotive components, cable insulation, and packaging materials.

Established blowing agents are exothermic blowing agents such as azodicarbonic acid diamide (azodicarbonamid, ADC, ADCA, CAS No. 123-77-3), the sulfonhydrazides 4,4'-oxybis(benzenesulfonylhydrazide) (OBSH, CAS No. 80-51-3) and p-toluenesulfonylhydrazide, (TSH, CAS No. 1576-35-8) and endothermic blowing agents such as carbonates, like sodium bicarbonate (SBC, NaHCO₃, CAS No. 144-55-8), and citric acid and its esters.

Since many years, azodicarbonamide (ADC) is one of the most effective and widely used chemical blowing agents for use in cellular thermoplastic and rubber applications (cf. for example DE-AS 1037 700). Azodicarbonamide decomposes on heating to give a high volume of gas, which consists mainly of nitrogen and carbon monoxide. These decomposition products are suitable to create a fine and uniform cell structured foam with a little shrink, a property which is fundamental in the production of soft foams such as plasticized PVC (P-PVC) or rubber foams. The decomposition temperature of azodicarbonamide can be reduced from 200 - 220 degrees centigrade to as low as 125 degrees centigrade by the addition of suitable activators (kickers), but useful decomposition rates are usually only achieved at 140 degrees centigrade and above. Activators or kickers are additives known in the art which are used to influence the decomposition temperature and rate of gas release of the blowing agent.

Azodicarbonamide may be combined with other chemical blowing agents in order to improve the processing behavior of the thermoplastic material and to optimize the end product. For example, in cellular rigid PVC (U-PVC; without softening of the polymer by adding plasticizers) applications such as foamed profile or sheet, ADCA may be used in combination with sodium bicarbonate to produce a foam structure with acceptable technical
performance (GB2314841). Because of differences in melt rheology, processing and demands on the foam structure, this technique cannot be transferred to plasticized, soft PVC and PVC plastisol processing.

However, azodicarbonamide as a blowing agent in plastics has been banned in the European Union since August 2005 for the manufacture of plastic articles that are intended to come into direct contact with food (COMMISSION DIRECTIVE 2004/1/EC of 6 January 2004 amending Directive 2002/72/EC as regards the suspension of the use of azodicarbonamide as blowing agent”. Official Journal of the European Union. 2004-01-13).

Furthermore, in December 2012, the European Chemicals Agency (ECHA) announced that azodicarbonamide was to be included in their Candidates List of Substances of Very High Concern (SY’HC) under Articles 57 and 59 of the Reach Regulation, which will limit or restrict the future use of ADCA. Therefore, there is a need for substitutes for ADCA having the same beneficial performance, especially for applications in foamed PVC.

If the most frequent used blowing agent, ADCA, is facing more and more challenges due to high concerns regarding safe and sustainable use of such additive in polymer formulations for example, then the demand for finding a suitable replacement to ADCA would be expected to increase. At any rate, for regions where the regulations are not as stringent, partial or total substitution of ADCA with a cost-effective, eco-friendly blowing agent alternative would provide an avenue for an end-user’s sustainability strategy and align with the overall trend to phase out such substance which is viewed as of high concern.

Possible alternative solutions are provided by the classes of sulfonylhydrazides and carbonates, but these substances exhibit some disadvantages when used as blowing agents, especially when used for applications in plasticized, soft PVC.

p-Toluenesulfonylhydrazide (TSH) starts decomposition at a temperature of about 105 degrees centigrade, which is considered as being too low for the processing of rigid and plasticised PVC. 4,4’-Oxybis(benzenesulfonylhydrazide) (OBSH) also releases nitrogen upon decomposition but the gas generation characteristic is different to that of azodicarbonamide. At temperatures above the decomposition point of OBSH the nitrogen release is rapid, but occurs at a different temperature compared to azodicarbonamide. Below the absolute product decomposition temperature of about 155 degrees centigrade, the decomposition and thus the gas release is slow. Additionally OBSH has the disadvantage that the decomposition products and the foamed end article produced have an unintended brownish discoloration at the typical P-PVC processing temperatures which are higher than 180 degrees centigrade.
The carbonates such as sodium bicarbonate do not liberate nitrogen but carbon dioxide and possibly water upon decomposition. Typically for carbon dioxide is its high solubility in the polymer, but it permeates out of the polymer matrix more rapidly than nitrogen, making it less efficient as a foaming agent, especially in plasticized PVC applications. The carbonates are generally not useful for the production of soft foams with a fine and uniform cell structure with little shrink. Sodium bicarbonate, the most common representative of the carbonates used as chemical blowing agents, has a slow decomposition and release of gas, which occurs over a wider temperature range in comparison to both ADCA and OBSH. The decomposition temperature of sodium bicarbonate can be influenced by citric acid.

Alkali metal bicarbonate particles, such as sodium bicarbonate particles and potassium bicarbonate particles, are known in the art. These products have many properties which make them interesting and extensively used in several technical fields, such as pharmaceutical industry, the feed and food industry, in detergents and in the treatment of non-ferrous metals.

The most common way to manufacture bicarbonate particles is crystallization by carbonization with carbon dioxide of a solution of the corresponding alkali metal (sodium or potassium carbonate for example) or a solution of the hydroxide of the corresponding alkali metal. It is also common to crystallize bicarbonates by controlled cooling of bicarbonate solutions, or by evaporating the solvent of such solutions.

For the industrial use of alkali metal bicarbonate particles control of specific properties of the particles, e.g., their bulk density (poured bulk density) or angle of repose is required. Some methods to control these parameters, such as a bulk density are known in the art. For example, US5411750 discloses a method of producing sodium bicarbonate powder with a bulk density between 70 and 500 kg/m³. The particles are prepared by spray-drying a dilute aqueous solution of the bicarbonate with an alkali metal salt as additive. WO 2014/096457 discloses a method for producing sodium bicarbonate particles by spray-drying of an aqueous solution comprising 1 - 10 % by weight of sodium bicarbonate and an additive selected from the group consisting of magnesium salt, sodium alkyl benzene sulfonate and soybean lecithin.

The above mentioned non-azodicarbonamide blowing agents fail to meet the expected requirement profile of a good blowing agent, and are in need of improvement in this respect.

SUMMARY

An aspect of the present invention provides a functionalized particulate bicarbonate, which can be used advantageously as a non-azodicarbonamide blowing agent in polymer
manufacture, particularly for the preparation of a foamed thermoplastic material, such as foamed PVC, polyurethanes, polyamides, polyolefins.

One aspect of the present invention relates to a functionalized particulate bicarbonate and its use as chemical blowing agent for foaming

- a thermoplastic polymer, or
- a polymer resin in an extrusion process.

The functionalized particulate bicarbonate may comprises ammonium bicarbonate, sodium bicarbonate and/or potassium bicarbonate, preferably sodium bicarbonate.

The functionalized particulate bicarbonate is functionalized with at least one additive selected from the group consisting of polyoxyalkynes, any derivatives thereof (such as ethers or esters), salts thereof, and any combination of two or more thereof. This additive may be called the "polyoxyalkylene additive".

The polyoxyalkylene additive may comprise or consist of:

- a polyethylene oxide (PEO) polymer, preferably of a MW of 20,000 or more,
- a polyethylene glycol (PEG) polymer, preferably of a MW of less than 20,000, or a MW of 4000 or less, or a MW of 3600 or less, or a MW of 2000 or less, or a MW of 1000 or less, or a MW of 800 or less,
- a polypropylene oxide (PPO) polymer,
- a polypropylene glycol (PPG) polymer,
- a co-polymer or block polymer which is made with at least one monomer of propylene oxide or propylene glycol,
- a co-polymer or block polymer which is made with at least one monomer of ethylene oxide or ethylene glycol,
- a polyethylene glycol / polypropylene glycol (PEG/PPG) block co-polymer, such as the PEG/PPG/PEG triblock polymer,
- any derivatives (such as ethers or esters) thereof,
- wherein any of these polymers may be chemically functionalized or not.

Polyethylene glycols are available in a wide range of different molecular weights. In one embodiment of the present invention low molecular weight polyethylene glycols having a molecular weight below 1000 g/mol can be used, preferably a polyethylene glycol having a molecular weight in the range of from 200 to 600 g/mol, such as in the range of from 300 to 500 g/mol, preferably PEG400. In another embodiment of the present invention a high molecular weight polyethylene glycol having a molecular weight of 1000 g/mol or above may be employed. Preferably the high molecular weight polyethylene glycol has a molecular
weight of from 1000 to 10000 g/mol, more preferably from 2000 to 8000 g/mol, such as PEG4000.

In some embodiments, the polyoxyalkylene additive is a polyoxyalkylene derivative which is an ester, such as a PEG or PPG ester. PEG or PPG esters may be polyethylene glycol or polypropylene glycol esters of fatty acids or resin acids.

Polyethylene glycols of high or low molecular weight (as defined above) and in particular low molecular weight polyethylene glycol and derivatives thereof, such as PEG or PPG esters are preferred polyoxyalkylene additives.

In some embodiments, the polyoxyalkylene additive is a polyoxyalkylene derivative which is an ether.

In some embodiments, the polyoxyalkylene additive is a polyoxyalkylene derivative which is a chemically functionalized polyoxyalkylene, for example, a polyoxyalkylene bearing one or more reactive substituents, such as -OH, -NH₂, -NHR, -SH and -COOH, wherein R is a hydrocarbon residue, and which has been reacted with a functional chemical compound being reactive with the reactive substituent of the polyoxyalkylene. Suitable functional chemical compounds are, for example, monofunctional or polyfunctional isocyanate or a monofunctional or polyfunctional epoxide.

In some embodiments, the polyoxyalkylene additive may comprise a refractive index of from 1.43 to 1.48, preferably a refractive index of from 1.44 to 1.47, more preferably a refractive index of from 1.45 to 1.46.

In some embodiments, the polyoxyalkylene additive may comprise a density measured at 20°C of from 1.0 to 1.2 g/ml, preferably a density measured at 20°C of from 1.05 to 1.18 g/ml, still more preferably a density measured at 20°C of from 1.08 to 1.16 g/ml.

In some embodiments, the polyoxyalkylene additive in the chemical blowing agent is in liquid form at 20°C;

In some embodiments, the polyoxyalkylene additive may be a defoamer. A "defoamer" also known as "anti-foaming agent" is a chemical additive that reduces and hinders the formation of foam in a liquid.

In some embodiments, the polyoxyalkylene additive may be an air release agent. An "air release agent" is designed to aid the de-airing of a foamable composition.

In some embodiments, the functionalized particulate bicarbonate may further comprise at least one additional additive (other than the polyoxyalkylene additive) selected from:
- one or more polymers (other than a polyoxyalkylene);
- one or more amino acids, any derivative thereof, and salts thereof;
- one or more inorganic salts;
- one or more oils;
- one or more fats;
- one or more resin acids, any derivative thereof, and salts thereof;
- one or more fatty acids, any derivative thereof, and salts thereof;
- a carboxylic or polycarboxylic acid, derivative thereof (such as esters), or salts thereof;
- one or more soaps;
- one or more waxes; or
- any combinations thereof;

preferably selected from the group consisting of:
- one or more fatty acids, any derivative thereof, and salts thereof;
- a carboxylic or polycarboxylic acid, derivative thereof (such as esters), or salts thereof;

and
- one or more resin acids, any derivative thereof, and salts thereof.

In some embodiments, the functionalized bicarbonate may further comprise silica. Silica may be used as a processing aid, an anti-caking agent and/or a flow aid for the functionalized bicarbonate. It is recommended that the silica is in amorphous (and noncrystalline) form. Preferably the silica in the functionalized bicarbonate is amorphous precipitated silica. The functionalized bicarbonate may comprise at least 0.1 wt%, preferably at least 0.2 wt%, more preferably at least 0.5 wt% of silica based on the total composition of the functionalized bicarbonate. It is recommended that the functionalized bicarbonate comprises not more than 5 wt% silica, preferably not more than 4 wt% silica, more preferably not more than 3 wt% silica.

The functionalized particulate bicarbonate may be spray-dried bicarbonate particles in the presence of the polyoxyalkylene additive, or co-milled bicarbonate particles in the presence of the polyoxyalkylene additive, or coated with the polyoxyalkylene additive in a fluid bed, or granulated with the polyoxyalkylene additive in a fluid bed, or coated with the polyoxyalkylene additive in an extruding device.

The functionalized particulate bicarbonate may comprises at least 50% by weight and less than 100% by weight of the bicarbonate component, and from 50% to 0.02% by weight of at least one additive (the polyoxyalkylene additive and optionally one or more additional additives). The functionalized particulate bicarbonate may comprise at least 65% by weight and less than 100% by weight of the bicarbonate component, and from 35% to 0.02% by weight of at least one additive (the polyoxyalkylene additive and optionally one or more
additional additives). The functionalized particulate bicarbonate may comprise at least 75% by weight and less than 100% by weight of the bicarbonate component, and from 25% to 0.02% by weight of at least one additive (the polyoxyalkylene additive and optionally one or more additional additives). The functionalized particulate bicarbonate may comprise at least 90% by weight and less than 100% by weight of the bicarbonate component, and from 10% to 0.02% by weight of at least one additive (the polyoxyalkylene additive and optionally one or more additional additives). Optionally, the functionalized bicarbonate may further comprise from 0.1 wt% to 5 wt% or from 0.2 wt% to 4 wt% or from 0.5 wt% to 3 wt% silica, preferably amorphous silica, more preferably amorphous precipitated silica.

Another aspect of the present invention relates to a chemical blowing agent for foaming a thermoplastic polymer, for example PVC plastisol or a polymer resin in an extrusion process, said chemical blowing agent comprising or consisting of the functionalized particulate bicarbonate, wherein said functionalized particulate bicarbonate contains at least the polyoxyalkylene additive.

In a preferred embodiment, the chemical blowing agent is endothermic.

In a preferred embodiment, the chemical blowing agent does not contain a blowing agent which is exothermic.

In a preferred embodiment, the chemical blowing agent does not contain a blowing agent which would liberate nitrogen gas and/or ammonia during heating when a foamed polymer is made using such chemical blowing agent.

In most preferred embodiments, the chemical blowing agent comprises or consists of the functionalized particulate sodium bicarbonate.

In some embodiments, the chemical blowing agent may further comprise a second compound which liberates CO2 upon heating, said second compound being selected from the group consisting of a carboxylic or polycarboxylic acid, derivative thereof (such as esters), or salts thereof. The second compound may comprise or may be at least one of:
- fumaric acid,
- tartaric acid,
- citric acid, citrates (such as sodium hydrogen citrate, disodium citrate), or esters of citric acid; or
- any combination thereof.

The second compound may be functionalized with at least one additive which is different or the same as the one used in the functionalized particulate bicarbonate. In some
embodiments, the second compound may be functionalized with at least the polyoxyalkylene additive.

Another aspect of the present invention relates to a foamable polymer composition comprising the functionalized particulate bicarbonate as blowing agent, wherein said functionalized particulate bicarbonate contains at least the polyoxyalkylene additive.

In some embodiments, the foamable polymer composition comprises the functionalized particulate bicarbonate as a first endothermic blowing agent and a carboxylic or polycarboxylic acid, ester thereof, or salt thereof, as a second endothermic blowing agent. The carboxylic or polycarboxylic acid, ester thereof, or salt thereof may be also functionalized. The functionalized particulate bicarbonate and the functionalized carboxylic or polycarboxylic acid, ester thereof, or salt thereof may be functionalized together or separately.

In particular embodiments, the foamable polymer composition does not contain a blowing agent which would liberate nitrogen gas and/or ammonia during heating when a foamed polymer is made from such foamable composition.

In some embodiments, the foamable polymer composition does not contain an exothermic blowing agent.

During foaming of polymer resin (plastisol or extrusion process) when particles of non-functionalized sodium bicarbonate are used as foaming agent, it was observed that the gas release occurs earlier than expected, due to the quick decomposition of bicarbonate. It has been found that functionalizing the bicarbonate particles by coating, by granulation, and/or by encapsulating with specific additives, in particular the polyoxyalkylene additive, enhances the protection of the bicarbonate particles with an inactive barrier, which delays the thermal decomposition when having various bicarbonate particle sizes, small (some may be nano-sized) and large (some may be micron-sized).

The functionalized particulate bicarbonate according to this aspect of the present invention is preferably a particulate sodium bicarbonate which is functionalized with at least one additive. This functionalized particulate sodium bicarbonate shows improved properties of expansion in comparison with nonfunctionalized particulate sodium bicarbonate of equivalent size. A “nonfunctionalized particulate sodium bicarbonate” is defined as a particulate sodium bicarbonate made without the additive(s) used in the making of the functionalized particulate sodium bicarbonate. The functionalized particulate sodium bicarbonate in the foamable polymer composition may reduce the time of gelation at a
temperature above glass transition temperature $T_g$ and/or above melting temperature $T_m$ of the polymer.

A process for manufacturing a polymer may comprise heating the foamable polymer composition which comprises the functionalized particulate bicarbonate at a temperature suitable for liberating $\text{CO}_2$ gas and melting the polymer during a gelation time at a temperature above glass transition temperature $T_g$ and/or above melting temperature $T_m$ of the polymer which is less than 130 seconds.

For some embodiments of a process for manufacturing a PVC polymer, when the foamable polymer composition is heated, the temperature suitable for liberating $\text{CO}_2$ gas from the functionalized particulate sodium bicarbonate and melting a PVC polymer may be from 190 to 210°C, preferably from 200 to 210°C, during a gelation time from 90 seconds to 120 seconds to provide a foamed PVC polymer.

When the foamable polymer composition is spread coated on a surface before heating and polymer melting, the foamed polymer may have an expansion ratio of at least 270, preferably at least 280, more preferably at least 300 and/or has a density of less than 0.6 g/cm$^3$, preferably less than 0.55 g/cm$^3$, more preferably at most 0.5 g/cm$^3$. The expansion ratio is calculated based on the ratio of a final thickness over an initial thickness of a spread coated layer of the foamable polymer composition as that layer gets heated during gelation in an oven.

The functionalized sodium bicarbonate particles are produced in the presence of at least one polyoxyalkylene additive from a solution containing sodium bicarbonate or directly from solid sodium bicarbonate particles.

The functionalized particulate bicarbonate can be obtained by at least one of the following processes:

- by spray-drying (also known as atomization), wherein the additive is dissolved in the bicarbonate-containing solution,
- by grinding or co-grinding (also known as milling or co-milling) with the additive(s) in emulsion or powder form,
- by spray coating and granulation within a fluidized bed,
- by spray agglomeration within a fluidized bed,
- by spray chilling (e.g., spray cooling, spray freezing),
- by roller compaction,
and/or
- by extrusion, including simultaneous mixing/extrusion.
It should be understood that one or more methods from this list may be omitted. In preferred embodiments, the functionalized particulate bicarbonate can be obtained by at least one of the following processes:
- by grinding or co-grinding (also known as milling or co-milling) with the additive(s) in emulsion or powder form;
- by spray coating and granulation within a fluidized bed, and/or
- by extrusion, including simultaneous mixing/extrusion.

In more preferred embodiments, the functionalized particulate bicarbonate can be obtained by at least one of the following processes:
- by spray coating and granulation within a fluidized bed, and/or
- by extrusion, including simultaneous mixing/extrusion.

In yet other preferred embodiments, the functionalized particulate bicarbonate can be obtained by at least one of the following processes:
- by grinding or co-grinding (also known as milling or co-milling) with the additive(s) in emulsion or powder form; and/or
- by extrusion, including simultaneous mixing/extrusion.

The functionalized particulate bicarbonate which is obtained by at least one said process may be further subjected to milling to reduce its mean particle size.

The process for functionalizing the particulate bicarbonate with any additional additive may be the same or different as the process used for functionalizing the same particulate bicarbonate with the polyoxyalkylene additive.

In some embodiments, the functionalized particulate bicarbonate may show excellent CO₂ release properties. The maximum loss temperature by TGA analysis of the functionalized particulate bicarbonate is preferably higher than the non-functionalized bicarbonate without the polyoxyalkylene additive. The CO₂ release of the functionalized particulate bicarbonate typically has its maximum at a temperature of at least 130 °C, preferably at a temperature of at least 135 °C, more preferably at a temperature of at least 140 °C, even more preferably at a temperature of at least 145 °C, and particularly preferably at a temperature of at least 155 °C.

In some embodiments, the maximum peak temperature determined by DSC thermal analysis, of the functionalized particulate bicarbonate is preferably higher than the non-
functionalized bicarbonate without the polyoxyalkylene additive. The DSC maximum peak temperature of the functionalized particulate bicarbonate with the polyoxyalkylene additive may be at least 140 °C, preferably at least 145 °C, more preferably at least 150 °C, even more preferably at least 155 °C, and particularly at least 160 °C.

5

DETAILED DESCRIPTION

Definitions

In the present description, wherein an element or composition is said to be included in and/or selected from a list of recited elements or components, it should be understood that in related embodiments explicitly contemplated here, the element or component can also be any one of the individual recited elements or components, or can also be selected from a group consisting of any two or more of the explicitly listed elements or components.

Further, it should be understood that elements and/or features of an apparatus, a process or method described herein can be combined in a variety of ways without departing from the scope and disclosures of the present teachings, whether explicit or implicit herein.

The term "thermoplastic material" shall mean a polymer that becomes pliable or moldable above a specific temperature, so is capable of flow at high temperatures below the thermal decomposition temperature and returns to a solid state upon cooling. A polymer is a macromolecular compound prepared by reacting (i.e. polymerizing, condensation) monomers of the same or different type, including homo- and copolymers. Thermoplastic materials are made by chain polymerization, polyaddition and / or polycondensation.

The term "functionalized particulate bicarbonate" is to be understood as defining particles which comprise a bicarbonate and an additive, preferably within the same particle. For example, the additive may form a layer or coating on the bicarbonate or the bicarbonate may form a layer or coating on the additive. Alternatively or additionally the additive may be embedded within a matrix of the bicarbonate or vice versa. The particle comprising bicarbonate and additive may be an agglomeration of smaller particles or small particles of one of the components may be agglomerated to a larger particle (or larger particles) of the other component. Preferably, the term "functionalized particulate bicarbonate" does not include a mere mixture of bicarbonate particles and at least one additive whether it be in liquid form or in the form of particles.

The term "functionalizing additive", as used herein, refers to a compound that is capable of improving at least one CO₂ release property of sodium bicarbonate when the additive is formulated with the sodium bicarbonate, relative to the sodium bicarbonate alone
(without additive). For example, the functionalizing additive is capable of increasing the CO₂ release beginning temperature and/or the CO₂ release maximum temperature of the functionalized particulate bicarbonate, determined in accordance with Example 54 of the present application. An example of a "functionalizing additive" in the present application is the polyoxyalkylene additive.

The term "comprising" includes "consisting essentially of" and "consisting of.

The term "foamed" in connection with the terms "thermoplastic material", "polymer" and "PVC" shall mean such material, polymer, or PVC with a cellular structure which is formed by gas generation from thermal decomposition and/or chemical reaction of a chemical blowing agent during processing.

The term "ppm" means parts per million, expressed by weight (e.g., 1 ppm = 1 mg / kg).

The term "per" means parts by weight of resin (e.g., 80 per of additive = 80 g additive per 100 g of resin).

The sign "%" or "wt%" refers to "weight percent" unless specifically stated otherwise.

The term "powder" shall mean a compound consisting of milled (ground), extruded, or spray-dried solid particles.

The term "exothermic blowing agent" defines a chemical that generates heat during its decomposition. An exothermic blowing agent typically undergoes rapid decomposition in a narrow temperature range. Generally speaking, the exothermic chemical blowing agents are associated as those chemicals giving N₂ as the main blowing gas (> 50 vol% of the generated gas is N₂). Other minor gases though may evolve from the decomposition of the exothermic chemical blowing agent. These other minor gases may include carbon monoxide, also in small amounts (< 5 vol%) of ammonia, and/or CO₂.

The term “endothermic blowing agent” defines a chemical that absorbs heat during its decomposition. An endothermic blowing agent typically has broader decomposition ranges in terms of both temperature and time. Most endothermic chemical blowing agents generate CO₂ as the main blowing gas (> 50 vol% of the generated gas is CO₂).

The CO₂ release properties of the functionalized particulate bicarbonate according to one aspect of the present invention can be determined by performing a thermogravimetric analysis (TGA) of a functionalized particulate bicarbonate sample, measuring the weight loss of the sample in dependence of the temperature. The CO₂ release properties are characterized by the derivate value for weight loss depending on the temperature. The CO₂ release beginning temperature is the temperature where the derivate value for weight loss starts to
raise. The C\textsubscript{0.2} release maximum temperature is the temperature where the derivate value for weight loss is at maximum. Typically, heating is performed between 30 °C and 250 °C at a speed of 10 °C/min. Thermogravimetric analysis can be performed for example on an STD Q600 V20.9 Build 20 thermogravimetric analysis instrument (provided by TA Instruments).

A plurality of elements includes two or more elements.

The phrase ‘A and/or B’ refers to the following selections: element A; or element B; or combination of elements A and B (A+B). The phrase ‘A and/or B’ is equivalent to at least one of A and B. The phrase ‘A and/or B’ equates to at least one of A and B.

The phrase A\textsubscript{1}, A\textsubscript{2}, ..., and/or A\textsubscript{n} with n \geq 3 includes the following choices: any single element A\textsubscript{i} (i= 1, 2, ... n); or any sub-combinations of from two to (n-1) elements chosen from A\textsubscript{1}, A\textsubscript{2}, ..., A\textsubscript{n}; or combination of all elements A\textsubscript{i} (i=T, 2, ... n). For example, the phrase A\textsubscript{1}, A\textsubscript{2}, and/or A\textsubscript{3} refers to the following choices: A\textsubscript{1}; A\textsubscript{2}; A\textsubscript{3}; A\textsubscript{1}+A\textsubscript{2}; A\textsubscript{1}+A\textsubscript{3}; A\textsubscript{2}+A\textsubscript{3}; or A\textsubscript{1}+A\textsubscript{2}+A\textsubscript{3}.

In the present specification, the description of a range of values for a variable, defined by a bottom limit, or a top limit, or by a bottom limit and a top limit, also comprises the embodiments in which the variable is chosen, respectively, within the value range: excluding the bottom limit, or excluding the top limit, or excluding the bottom and the top limit.

In the present specification, the description of several successive ranges of values for the same variable also comprises the description of embodiments where the variable is chosen in any other intermediate range included in the successive ranges. Thus, for illustration purpose, when it is stated that "the element X is generally at least 10, advantageously at least 15", the present description also includes another embodiment where a new minimum can be selected between 10 and 15, for example: where "the element X is at least 11", or also where: "the element X is at least 13.74", etc.; 11 or 13.74 being values included between 10 and 15. Also for illustration purpose, when it is indicated that "the element X is generally at most 15, advantageously at most 10", the present description also includes another embodiment where a new maximum can be selected between 10 and 15.

In the present description, wherein an element or composition is said to be included in and/or selected from a list of recited elements or components, it should be understood that in related embodiments explicitly contemplated here, the element or component can also be any one of the individual recited elements or components, or can also be selected from a group consisting of any two or more of the explicitly listed elements or components.

For example, when in an embodiment the choice of an element from a group of elements is described, the following embodiments are also explicitly described:
- the choice of two or more elements from the group,
- the choice of an element from a subgroup of elements consisting of the group of elements from which one or more elements have been removed.

The use of the singular 'a' or 'one' herein includes the plural unless specifically stated otherwise.

In addition, if the term "about" or "ca." is used before a quantitative value, the present teachings also include the specific quantitative value itself, unless specifically stated otherwise. As used herein, the term "about" or "ca." refers to a +10% variation from the nominal value unless specifically stated otherwise.

FUNCTIONALIZED PARTICULATE BICARBONATE

One aspect of the present invention relates to a functionalized particulate bicarbonate. The functionalized particulate bicarbonate comprises a bicarbonate ingredient which preferably is an alkali or ammonium salt, such as sodium bicarbonate, potassium bicarbonate, and/or ammonium bicarbonate, sodium and potassium bicarbonate being preferred, in particular sodium bicarbonate being most preferred.

In some embodiments, the functionalized particulate bicarbonate preferably comprises at least 50% by weight, or at least 55% by weight, or at least 60% by weight, or even at least 65% by weight, but less than 100% by weight of the bicarbonate ingredient (e.g., ammonium, sodium and/or potassium bicarbonate).

In some embodiments, the functionalized particulate bicarbonate may comprise at least 90% by weight, or at least 93% by weight, or at least 94% by weight, or even at least 95% by weight, but less than 100% by weight of the bicarbonate ingredient (e.g., ammonium, sodium and/or potassium bicarbonate).

In some particular embodiments, the functionalized particulate bicarbonate preferably comprises at least 90% by weight but less than 100% by weight of an alkali metal bicarbonate. The functionalized particulate bicarbonate preferably comprises at least 92% by weight of an alkali metal bicarbonate, at least 93% by weight, more preferably at least 94% by weight, in particular at least 95% by weight of the alkali metal bicarbonate, in particular sodium bicarbonate.

In some embodiments, the functionalized particulate bicarbonate may have 50% by weight or less, or 45% by weight or less, or 40% by weight or less, or even 35% by weight or less, of at least one functionalization additive (the polyoxyalkylene and optionally one or more additional functionalization additives).
In some particular embodiments, the functionalized particulate bicarbonate contains 10% by weight or less, or 7% by weight or less, or 5% by weight or less, or 3% by weight or less, of at least one functionalization additive (the polyoxyalkylene additive and optionally one or more additional functionalization additives).

The functionalization additive (the polyoxyalkylene and optionally one or more additional functionalization additives) should be present in the functionalized particulate bicarbonate according to the present invention in the amount of at least 0.02% by weight, preferably at least 0.05% by weight, in particular at least 0.1% by weight. The higher the weight% of the functionalization additive in the functionalized particulate bicarbonate, the more disadvantageous it may be for cost reasons. Preferably, to reduce cost of the more expensive functionalization additive compared to the bicarbonate ingredient, it may be desirable to use at most 8% by weight, more preferably at most 6% by weight, in particular at most 5% by weight of the additive in the functionalized particulate bicarbonate, wherein the functionalization additive comprises or consists of the polyoxyalkylene additive.

However in embodiments where the functionalization additive is relatively not expensive (for example when its cost is not more than twice that of the bicarbonate ingredient), it may be desirable to use at least 5% by weight, more preferably at least 7% by weight, in particular at least 10% by weight of the functionalization additive and/or at most 50% by weight, more preferably at most 40% by weight, yet more preferably at most 35% by weight of the functionalization additive in the functionalized particulate bicarbonate, wherein the functionalization additive comprises or consists of the polyoxyalkylene additive.

In some particular embodiments, the functionalized particulate bicarbonate may comprise 0.02-50%, or 0.02-45%, or 0.02-40%, or 0.02-35%, by weight of at least one functionalization additive (the polyoxyalkylene additive and optionally one or more additional functionalization additives).

In some particular embodiments, the functionalized particulate bicarbonate may comprise more than 1% and up to 50% by weight of the polyoxyalkylene additive.

In some embodiments, for cost effectiveness, the functionalized particulate bicarbonate may comprise from 1% to 25% by weight, preferably from 3% to 20% by weight, of the polyoxyalkylene additive.

In particular embodiments, the functionalized particulate bicarbonate may comprise at least 65% by weight and less than 100% by weight of the bicarbonate component, and from 35% to 0.02% by weight of at least one additive; or may comprise at least 75% by weight and
less than 100% by weight of the bicarbonate component, and from 25% to 0.02% by weight of at least one additive.

The functionalized particulate bicarbonate is preferably used as a blowing agent for foamed or extruded polymers (such as foamed PVC or polyurethanes: extruded PVC, polyolefns, polyamides), preferably used as an endothermic blowing agent. Non-limiting examples of polymers are polyvinyl chloride (PVC), polyurethanes, polyolefns (PO, PE, PP), styrenics (PS, ABS, ASA, SAN) and natural and synthetic rubber such as nitrile butadiene rubber (NBR) or chloroprene rubber (CR), polyamides, polyimides.

The functionalized particulate bicarbonate may further contain an additive which is capable of liberating C02 and which is also used for functionalizing the particulate bicarbonate. This additive may be considered as a secondary blowing agent in the functionalized particulate bicarbonate. Not only this additive would provide an increase in the C02 generation when the functionalized particulate bicarbonate is used as an endothermic blowing agent, but also this additive would protect the bicarbonate core from premature C02 release by protecting its surface (or part thereof). This C02-liberating additive may be a carboxylic or polycarboxylic acid, derivative thereof (such as esters), or salts thereof.

Suitable carboxylic acids include those of the formula: HOOC--R—COOH where R is an alkylene group of 1 to about 8 carbon atoms which may also be substituted by one or more hydroxy groups or keto groups and may also contain unsaturation. Also included are esters, salts and half salts.

A preferred C02-liberating additive may include at least one of:
- fumaric acid,
- tartaric acid, or
- citric acid, citrates (such as sodium hydrogen citrate, disodium citrate), or esters of citric acid.

Esters of citric acid may include tributyl citrate, triethyl citrate, tri-C12-13 alkyl citrate, tri-C14-15 alkyl citrate, tricaprylyl citrate, triethyhexyl citrate, triisocetyl citrate, trioctylidodecyl citrate and triisostearyl citrate, isodecyl citrate and stearyl citrate, dilauryl citrate, and/or ethyl citrates (mixture of tri-, di- and monoesters), preferably tributyl citrate, triethyl citrate, isodecyl citrate, or triethyhexyl citrate.

A more preferred C02-liberating additive comprises or consists of citric acid, esters thereof, or salts thereof.
In some embodiments, the functionalized particulate bicarbonate does not contain citric acid, esters thereof, or salts thereof.

In some embodiments, the functionalized particulate bicarbonate does not contain an exothermic blowing agent.

In some particular embodiments, the functionalized particulate bicarbonate does not contain a compound used as blowing agent which liberates ammonia.

In some particular embodiments, the functionalized particulate bicarbonate does not contain a compound used as blowing agent which liberates nitrogen gas. Examples of blowing agents which liberate nitrogen gas are exothermic blowing agents such as azodicarbonic acid diamide (azodicarbonamide, ADC, ADCA, CAS No. 123-77-3), the sulfonhydrazides 4,4′-oxybis(benzenesulfonylhydrazide) (OBSH, CAS No. 80-51-3) and p-toluenesulfonylhydrazide, (TSH, CAS No. 1576-35-8).

In preferred embodiments, the functionalized particulate bicarbonate does not contain azodicarbonamide.

In alternate or additional preferred embodiments, the functionalized particulate bicarbonate does not contain benzenesulfonylhydrazide.

In alternate or additional embodiments, the functionalized particulate bicarbonate does not contain p-toluenesulfonylhydrazide.

In preferred embodiments of the present invention, the functionalized particulate bicarbonate comprises the bicarbonate ingredient and the at least one additive in powder form.

For certain applications it is preferred that the functionalized particulate bicarbonate of the present invention contains sodium bicarbonate as particles which are coated with a coating of the functionalization additive. Such coating can improve some properties of the functionalized particulate bicarbonate. The functionalization additive in such instance may be termed a "coating agent". The additive as coating agent shall mean that this additive is able to cover, partly or completely, the surface of particles of bicarbonate. The "coating agent" is a different compound than bicarbonate ingredient from which the core of the particles are made of.

For certain applications, it is envisioned that the functionalized particulate bicarbonate of the present invention contains sodium bicarbonate co-milled with one functionalization additive. Such co-milling with the functionalization additive can improve some properties of the functionalized particulate bicarbonate.
For certain applications, the functionalized particulate bicarbonate of the present invention contains sodium bicarbonate as particles which are functionalized with more than one functionalization additive, so long as one functionalization additive is a polyoxyalkylene additive. The functionalization of the sodium bicarbonate particles may be carried out simultaneously with the additives using one functionalization method, or may be carried out sequentially using one additive with one functionalization method and then another additive with the same or different functionalization method. For example, sodium bicarbonate particles may be first functionalized with a first additive, and then these first-functionalized particles are again functionalized with a second additive (the second additive having the same composition or a different composition as the first additive, preferably having a different composition). The methods used for subsequent functionalizations may be the same, but preferably are different. The (first and second) functionalization methods are preferably selected from the group consisting of extrusion, co-grinding, and spray coating. For example, the first functionalization method may comprise or consist of co-grinding or extrusion, and the second functionalization method may comprise or consist of extrusion, co-grinding, or spray coating. Preferably, the first functionalization method may comprise or consist of co-grinding, and the second functionalization method may comprise or consist of extrusion. At least one of the first and second functionalization additives is the polyoxyalkylene additive. Preferably the second functionalization additive is the polyoxyalkylene additive, and the second functionalization method may comprise or consist of extrusion.

For certain applications, the functionalized particulate bicarbonate of the present invention contains sodium bicarbonate as particles which are functionalized with the polyoxyalkylene additive, but the polyoxyalkylene additive is not added all at once, but added sequentially in several portions. For example, bicarbonate particles may be first functionalized with a first portion of the polyoxyalkylene additive, and then these first-functionalized bicarbonate particles are again functionalized with a second portion of the same polyoxyalkylene additive. The methods used for functionalization may be the same or may be different. For example, the (first and second) functionalization methods are preferably selected from the group consisting of extrusion, co-grinding, and spray coating. Preferably, the first functionalization method may comprise or consist of co-grinding, and the second functionalization method may comprise or consist of extrusion.

For certain applications, it may be preferred that the functionalized particulate bicarbonate of the present invention contains sodium bicarbonate as particles which are coated with a coating of a first additive, and then these coated particles are co-milled with a
second additive (the second additive having the same composition or a different composition as the first additive).

For certain applications, the production of a fine cellular foam, so to produce a certain small amount of gas at one place, might be desirable. In order to improve the cellular foam structure it can be suitable that the functionalized particulate bicarbonate has a characteristic particle size and particle size distribution. The $D_{50}$ term is designating the diameter for which 50% by weight of the particles have a diameter less than or equal to $D_{50}$ (weight average diameter). The $D_{10}$ term is designating the diameter for which 10% by weight of the particles have a diameter less than or equal to $D_{10}$. The $D_{90}$ term is designating the diameter for which 90% by weight of the particles have a diameter less or equal to $D_{90}$.

The functionalized particulate bicarbonate may have advantageous properties, such as a low particle size preferably with a low span. The span of the particle size distribution is as known in the art defined as the ratio $(D_{90} - D_{10}) / D_{50}$. The span may range from about 1 to about 6, such as from about 1 to about 3. In one embodiment the span may be lower than 6, preferably lower than 4, more preferably lower than 3. In one embodiment the span may be higher than 1, preferably higher than 2. In another embodiment the span may be lower than 1.8, more preferably at most 1.7 in particular at most 1.6, e.g. at most 1.5.

Preferably, the particles of the functionalized particulate bicarbonate have a particle size distribution of $D_{50}$ of at most 250 µm, preferably at most 100 µm, more preferably at most 60 µm, yet more preferably at most 40 µm, or at most 30 µm, or at most 25 µm.

In some embodiments, the particles of the functionalized particulate bicarbonate have a particle size distribution of $D_{50}$ of more than 1 µm, preferably more than 2 µm, more preferably more than 5 µm, yet more preferably at least 8 µm. This functionalized particulate bicarbonate is termed "functionalized micron-sized bicarbonate".

In some embodiments, the particles of the functionalized particulate bicarbonate have a $D_{90}$ in the range of 1 µm - 160 µm, preferably in the range of 1 µm - 10 µm, more preferably in the range of 2 µm - 10 µm, yet more preferably in the range of 4 µm - 8 µm, in particular 5 µm - 6 µm.

In some embodiments, the particles of the functionalized particulate bicarbonate have a $D_{90}$ in the range of from 20 µm to 450 µm, preferably from 30 µm to 200 µm, more preferably from 30 µm to 165 µm, in particular from 30 µm to 100 µm.

The weight-average diameter $D_{50}$, as well as $D_{90}$ and $D_{90}$ values can be measured by laser diffraction and scattering on a Malvern Mastersizer S particle size analyser using an He-Ne laser source having a wavelength of 632.8 nm and a diameter of 18 mm, a measurement
cell equipped with a backscatter 300 mm lens (300 RF), and MS 17 liquid preparation unit, and an automatic solvent filtration kit ("ethanol kit") using ethanol saturated with bicarbonate (wet method).

In preferred embodiments, the functionalized particulate bicarbonate shows excellent CO₂ release properties. The maximum loss temperature determined by TGA analysis of the functionalized particulate bicarbonate is preferably higher than the non-functionalized bicarbonate without the polyoxyalkylene additive. The CO₂ release of the functionalized particulate bicarbonate may have its maximum at a temperature of at least 130 °C, preferably at a temperature of at least 135 °C, more preferably at a temperature of at least 140 °C, even more preferably at a temperature of at least 145 °C, and particularly at a temperature of at least 155 °C.

As determined by Differential Scanning Calorimetry (DSC) thermal analysis, the functionalized particulate bicarbonate preferably has the maximum peak temperature higher than the non-functionalized bicarbonate without the polyoxyalkylene additive. The DSC maximum peak temperature of the functionalized particulate bicarbonate may be at least 140 °C, preferably at least 145 °C, more preferably at least 150 °C, even more preferably at a temperature of at least 155 °C, and particularly preferably at a temperature of at least 160 °C.

FUNCTIONALIZED NANO-SIZED BICARBONATE

In some embodiments, the particles of the functionalized particulate bicarbonate have a particle size distribution of D₅₀ of at most 1 µm, preferably less than 1 µm. This functionalized particulate bicarbonate is termed "functionalized nano-sized bicarbonate".

In the instance when the functionalized particulate bicarbonate is based on nano-sized bicarbonate particles, it is preferred that the bicarbonate nano-sized particles are formed before functionalization. Techniques such as wet grinding with a solvent, micronisation and dry nanogrinding would be effective. Use of mills such as tumbler ball mills, planetary ball mills (for example available from Retch) or jet mills (for example available from Alpine) is suitable for making nano-sized bicarbonate particles. Ball milling involves decomposition of bulk solid materials into nanoscale regimes using a mechanical force. The reduction of particle size by high energy ball milling is termed as mechanical milling. Since milling a powder of bicarbonate powder to nanosize level generates quite a lot of heat, it is recommended to cool during the milling. Additionally, to facilitate the milling to nanosize level, it may be recommended to use a lubricant.
Additionally, to prevent particles from re-agglomerating during milling or after exiting the mill, it may be recommended to use a surfactant. These nano-sized particles have strong tendency to agglomerate owing to having large specific surface area. Surfactants may play an important role to prevent this close contact of the nano-sized particles by providing steric barrier and reducing surface tension. The surfactant molecules form a thin organic layer around the newly formed surfaces to protect the exposed surface from cold yielding when they come into contact with another surface during the milling process or when they exit the mill. A suitable surfactant may include a polymer such as poly(acrylic acid, sodium salt), or a fatty acid or ester thereof, such as oleic acid, stearic acid, oleic acid or oleylamine, palmitic acid, mystic acid, undecanoic acid, octanoic acid, and/or valeric acid.

Because the functionalization adds another compound (preferably the polyoxyalkylene additive) to the nanosized bicarbonate core particles, it is recommended to select a technique for additive deposition/incorporation (techniques being described in more detail below) which would not significantly increase the size of the initial bicarbonate core particles. It would be preferred for example for the particulate bicarbonate initially having a D50 of 1 μm or less before functionalization to generate after functionalization a functionalized particulate bicarbonate still in the nanosized range with a D50 of at most 1 μm or less. It is though acceptable in some instances that the functionalized particulate bicarbonate starting from nanosized bicarbonate core particles may reach a D50 of 2 μm or less after functionalization.

**ADDITIVE IN FUNCTIONALIZED PARTICULATE BICARBONATE**

The functionalization additive in the functionalized particulate bicarbonate comprises or consists of the polyoxyalkylene additive.

Polyoxyalkylene additives are known as defoamers and air release additives in the manufacture of foamed polymer compositions. They are incorporated for example into PVC plastisols to prevent the formation of foam and bubbles during production. The present inventors found that in combination with particulate bicarbonate, polyoxyalkylene additives exhibit a further function by increasing the foaming effectivity of the chemical blowing agent. This finding was particularly surprising because defoamers and air release additives are usually used to destabilize foam bubbles and to accelerate the migration of bubbles to the surface of a polymer film.

Suitable polyoxyalkylene additives can be selected by a person skilled in the art according to the requirements. For example, the polyoxyalkylene additive should be compatible with the polymer which is intended to be foamed using the chemical blowing
agent of the invention. Compatibility with the polymer depends for example on the molecular weight of the polyoxyalkylene additive and, if present, chemical substituents attached to the polyoxyalkylene additive.

In the context of the present invention, the polyoxyalkylene additive may comprise or consist of:

- a polyethylene oxide (PEO) polymer, preferably of a MW of 20,000 or more,
- a polyethylene glycol (PEG) polymer, preferably of a MW of less than 20,000, or a MW of 4000 or less, or a MW of 3600 or less, or a MW of 2000 or less, or a MW of 1000 or less, or a MW of 800 or less,
- a polypropylene oxide (PPO) polymer,
- a polypropylene glycol (PPG) polymer,
- a co-polymer or block polymer which is made with at least one monomer of propylene oxide or propylene glycol.
- a co-polymer or block polymer which is made with at least one monomer of ethylene oxide or ethylene glycol,
- a polyethylene glycol / polypropylene glycol (PEG/PPG) block co-polymer, such as the PEG/PPG/PEG triblock polymer,
- any derivatives (such as esters) thereof,
- wherein any of these polymers may be chemically functionalized or not.

Polyethylene glycols are available in a wide range of different molecular weights. In one embodiment of the present invention low molecular weight polyethylene glycols having a molecular weight below 1000 g/mol can be used, preferably a polyethylene glycol having a molecular weight in the range of from 200 to 600 g/mol, such as in the range of from 300 to 500 g/mol, preferably PEG400. In another embodiment of the present invention a high molecular weight polyethylene glycol having a molecular weight of 1000 g/mol or above may be employed. Preferably the high molecular weight polyethylene glycol has a molecular weight of from 1000 to 10000 g/mol, more preferably from 2000 to 8000 g/mol, such as PEG4000.

In some embodiments, the polyoxyalkylene additive is a polyoxyalkylene derivative which is an ester, such as a PEG or PPG ester. PEG or PPG esters which may be suitable as polyoxyalkylene additive may be polyethylene glycol or polypropylene glycol esters of fatty acids or resin acids.
Polyethylene glycols of high or low molecular weight (as defined above) and in particular low molecular weight polyethylene glycol and derivatives thereof, such as PEG or PPG esters are preferred polyoxyalkylene additives.

Suitable examples of PEG esters may be PEG monolaurate, PEG dilaurate, PEG dioctoate, PEG monooleate, PEG dioleate, PEG monostearate, PEG distearate, PEG monotallate, and/or PEG ditallate. These PEG esters come in a variety of molecular weights. For example, PEG monolaurate may be found as PEG-2 Laurate, PEG-2 Laurate SE, PEG-4 Laurate, PEG-6 Laurate, PEG-8 Laurate, PEG-9 Laurate, PEG-10 Laurate, PEG-12 Laurate, PEG-14 Laurate, PEG-20 Laurate, PEG-32 Laurate, PEG-75 Laurate, PEG-150 Laurate. A molecular weight of less than 1000 is generally preferred for a PEG ester used as a polyoxyalkylene additive. Some PEG esters can be found under the brandname Pegosperse® from Lonza and the brandname Lumulse from Vantage. Other suppliers are Americam eChem Inc. and Ivanhoe Industries, Inc.

Suitable examples of propylene glycol esters may be PPG stearate, PPG oleate, PPG laurate, or PPG myristate.

Polyoxyalkylene defoamers and air release additives are also commercially available for example from BYK-Chemie GmbH, Germany. A suitable and in the present invention particularly preferred polyoxyalkylene additive is the BYK-3155 silicone-free polymer-based air release additive obtainable from BYK-Chemie GmbH.

In some embodiments, the polyoxyalkylene additive is a polyoxyalkylene derivative which is a chemically functionalized polyoxyalkylene, for example, a polyoxyalkylene bearing one or more reactive substituents, such as -OH, -NH₂, -NHR, -SH and -COOH, wherein R is a hydrocarbon residue, and which has been reacted with a functional chemical compound being reactive with the reactive substituent of the polyoxyalkylene. Suitable functional chemical compounds are, for example, monofunctional or polyfunctional isocyanate or a monofunctional or polyfunctional epoxide.

The hydrocarbon residue in the above reactive substituent -NHR can be virtually any substituent comprising carbon and hydrogen atoms. The hydrocarbon can comprise alkyl, cycloalkyl, aryl and aralkyl and heteroatoms, such as O, N and S. The hydrocarbon can be saturated or unsaturated.

As chemically functionalized polyoxyalkylene additive, a polymer obtainable from the reaction of

A) polyepoxides with

B) at least one primary polyoxyalkyleneamine of the general formula (1)
where \( R = \text{C}1-\text{C}4 \) alkyl, \( \text{C}5-\text{C}12 \) cyeloalkyl, \( \text{C}6-\text{C}18 \) aryl, \( \text{C}6-\text{C}18 \) aralkyl, \( R^1 \) and \( R^2 \) independently of one another may be identical or different and are \( \text{H} \), \( \text{C}1-\text{C}4 \) alkyl, aryl, and/or \(-\text{CH}2-\text{O}-\text{C}6-\text{H}2+1\), and \( x \) may be arranged blockwise or randomly.

C) at least one other aliphatic and/or araliphatic primary amine of the general formula

\[
(\text{II}) \quad \text{H}2\text{N}-R^6-Z
\]

where \( R^6=\text{alkyl, aryl and aralkyl} \) and \( Z \) may be a further functional group selected from \(-\text{OH}, \) tertiary amine or a heterocyclic radical with a 5- or 6-membered ring.

D) at least one modified isocyanate of the general formula (IIia) or (IIlb)

\[
(\text{IIia}) \quad \text{R}^3-\text{O}-[\text{O}-\text{C} \text{Y} \text{O}]_{\text{m}} [\text{\text{C}CH} \text{CH} \text{O}]-_{\text{n}} \text{C} \text{N} \text{H}- \text{X}-\text{NCO}
\]

\[
(\text{IIlb}) \quad \text{R}^3-\text{O}-[\text{\text{C}CH} \text{CH} \text{O}]-_{\text{n}} [\text{O}-\text{C} \text{Y} \text{O}]_{\text{m}} \text{C} \text{N} \text{H}- \text{X}-\text{NCO}
\]

where \( \text{R}^3=\text{alkyl, cyeloalkyl, aryl and/or aralkyl} \), \( \text{R}^4 \) and \( \text{R}^5 \) independently of one another are \( \text{H} \), alkyl and/or aryl, \( \text{X}=\text{alkylene, cycloalkylene and/or aralkylene} \), \( \text{Y}=\text{alkylene and/or cycloalkylene} \), and \( \text{n and m} \) independently of one another are 0 to 100, where the sum \( n+m>2 \),

to form a urethane can be exemplified. This polymer is described in US 2013/0018126 A1, the content of which is incorporated herein by reference.

Another suitable chemically functionalized polyoxyalkylene additive is a biuret compound of the following general formula
This compound is disclosed in US 2007/0225451 Al, the content of which is incorporated herein by reference.

Another suitable chemically functionalized polyoxyalkylene additive is an addition compound produced by a process of reacting

a) one or more polyisocyanates containing uretdione groups with

b) one or more compounds of the formula (I)

\[ Y-(XH)_n \]

where

\( Y \) is a group that is reactive towards isocyanates and

\( XH \) is a group that is reactive towards isocyanates and

\( n \) is 1, 2 or 3, and

and when more than one compound of formula (I) is used, for at least 50 mol% of the compounds of formula (I), \( n \) shall equal 1,

with the proviso that at least 50% of the free isocyanate groups of the polyisocyanate, component a), are reacted with the compounds of the formula (I) to give an intermediate which contains uretdione groups,
and subsequently the intermediate is reacted with
c) one or more compounds of the general formula (II)
\[ \text{Z-}(\text{NHR})_m \]
in which
\[ \text{R} \] is hydrogen, an aliphatic, cycloaliphatic, araliphatic and/or aromatic radical having up to 36 C atoms, or a polyoxyalkylene radical,
\[ m \] is 1, 2 or 3 and
\[ \text{Z} \] is a non-basic aliphatic, cycloaliphatic, araliphatic and/or aromatic radical and/or polyoxyalkylene or polisiloxane radical having a number-average molar mass of less than 10,000 g/mol, which if desired may contain the following functional groups:

- \(-\text{OH}\)
- \(-\text{COOH}\)
- \(-\text{SO}_3\text{H}\)
- \(-\text{P}_3\text{H}_2\)
- \(-\text{Si(OR)}_3\) and \(-\text{SiR(OR)}_2\)

where the hydroxy[ groups are reacted if desired with polyphosphonic acid or \(\text{P}_2\text{O}_5\) or with polycarboxylic acids having at least two carboxyl groups or their anhydrides in an amount such that for each OH group to be reacted there is at least 0.8 molecule of polycarboxylic acid or polycarboxylic anhydride employed;

with the proviso that all free NCO groups that may still be present and at least 20 % of the uretdione groups used originally are reacted with \(\text{Z-}(\text{NHR})_m\). These addition compounds are disclosed in US 2007/0259120 A1, the content of which is incorporated herein by reference.

Another suitable chemically modified polyoxyalkylene additive can be obtained by reacting:

(I) monofunctional or polyfunctional aromatic epoxides; with

(II) polyoxyalkylenemonoamines having a number-average molecular weight of >400 g/mol, one primary or secondary amino group and at least 4 ether oxygen atoms per molecule, wherein

from 90 to 100 % of the epoxide groups of the starting material are reacted,

the weight fraction of aromatic groups in the addition compounds is less than 50 % and

the addition compounds containing per molecule at least one amino group on which salts can be formed, or being present as a salt. These polymers are disclosed in US 2005/0020735 A1, the content of which is incorporated herein by reference.
All preferred embodiments, such as preferred amounts of the polyoxyalkylene additive in the functionalized particulate bicarbonate are those as described below with respect to the optional additional additives.

The functionalization additive in the functionalized particulate bicarbonate may additionally comprise, as non-limiting examples, at least one following compound (additional additive):
- one or more polymers;
- one or more amino acids, any derivative thereof, and salts thereof;
- one or more inorganic salts;
- one or more oils;
- one or more fats;
- one or more resin acids, any derivative thereof, and salts thereof;
- one or more fatty acids, any derivative thereof, and salts thereof;
- a carboxylic or polycarboxylic acid, derivative thereof (such as esters), or salts thereof;
- one or more soaps;
- one or more waxes; or
- any combinations thereof.

In some embodiments, the additional additive may comprise or consist of a polymer selected from the group consisting of polyvinylalcohol, polyglycol, polysaccharide, poly(meth)acrylic acid, poly(acrylic acid co-maleic acid, polyethylenenimine, polyvinylpyrrolidone, N-2(-Hydroxypropyl) methacrylamide, and combinations thereof.

In some embodiments, the additional additive may comprise or consist of a polysaccharide selected from the group consisting of hydrolyzed starch, carboxymethylcellulose, alginic acid and its salt, arabic gum, carrageenan; guar gum, locust bean gum, xantham gum and combinations thereof.

In some embodiments, the additional additive may comprise or consist of an amino acid, derivative thereof or salt thereof selected from the group consisting of casein, gelatin, glycine, proline, hydroxyproline, glutamic acid, alanine, arginine, aspartic acid, lysine, pectin, serine, leucine, valine, phenylalanine, threonine, isoleucine, hydroxylysine, methionine, histidine, tyrosine and combinations thereof.

In some embodiments, the additional additive may comprise or consist of an inorganic salt selected from the group consisting of silicates (e.g., sodium silicate), NaCl, KCl, MgCl2, sodium phosphate, borates, nitrates, nitrites, sulfates, sulfites and combinations thereof.
In some embodiments, the additional additive may comprise or consist of:

- an amino acid, derivative thereof, or salt thereof,
- a polysaccharide (such as hydrolyzed starch, gums, carboxymethylcellulose),
- a resin acid, derivative thereof, or salt thereof,
- a fatty-acid, derivative thereof (such as esters), or salt thereof,
- a carboxylic or polycarboxylic acid, derivative thereof (such as esters), or salts thereof; or
- any combination thereof.

In some embodiments, the additional additive may comprise or consist of:

- a polymer other than the polyoxyalkylene additive (such as poly(meth)acrylates and derivatives thereof, polyvinylalcohol and polysaccharides, including modified, in particular hydrolyzed starch, maltodextrin and gum arabic),
- an amino acid, derivative thereof, or salt thereof (such as leucine),
- an oil (such as epoxidized soy bean oil),
- a resin acid, derivatives thereof, or salt thereof (such as rosin acid),
- a fatty acid, derivatives thereof, or salt thereof (such as stearic acid, lauric acid, linoleic acid and glycerol mono stearate),
- a wax (such as bees wax and carnauba wax), or
- any combination thereof.

In some embodiments, the additional additive may comprise or consist of a compound which is capable of liberating CO2 and which is also used for functionalizing the particulate bicarbonate. This additive may be considered as a secondary blowing agent in the functionalized particulate bicarbonate. Not only this additive would provide an increase in the CO2 generation when the functionalized particulate bicarbonate is used as an endothermic blowing agent, but also this additive would protect the bicarbonate core from premature CO2 release by protecting its surface (or part thereof). This CO2-liberating additive may be a carboxylic or polycarboxylic acid, derivative thereof (such as esters), or salts thereof.

Suitable carboxylic acids include those of the formula: HOOC—R—COOH where R is an alkylene group of 1 to about 8 carbon atoms which may also be substituted by one or more hydroxy groups or keto groups and may also contain unsaturation. Also included are esters, salts and half salts.
A preferred C02-liberating additive may include at least one of:

- fumaric acid,
- tartaric acid, or
- citric acid, citrates (such as sodium hydrogen citrate, disodium citrate), or esters of citric acid.

Esters of citric acid may include tributyl citrate, triethyl citrate, tri-C12-13 alkyl citrate, tri-C14-15 alkyl citrate, tricaprylyl citrate, triethyhexyl citrate, triisocetyl citrate, trioctyldodecyl citrate and tristearyl citrate, isodecyl citrate and stearyl citrate, dilauryl citrate, and/or ethyl citrates (mixture of tri-, di- and monoesters), preferably tributyl citrate, triethyl citrate, isodecyl citrate, or triethylhexyl citrate.

A more preferred C02-liberating additive comprises or consists of citric acid, any ester thereof, or any salt thereof.

In some embodiments, the additive excludes citric acid, esters thereof, or salts thereof.

In some embodiments, the additional additive may comprise or may consist of an amino acid, a derivative thereof, or a salt thereof.

Generally, the amino acids are compounds as known in the art composed of an amino group and a carboxylic acid functional group. An amino group is, in accordance with IUPAC nomenclature a compound formally derived from ammonia (N\(^3\)) by replacing one, two or three hydrogen atoms by hydrocarbyl groups, and having the general structures RNH\(_2\) (primary amines), R\(_2\)NH (secondary amines) or R\(_3\)N (tertiary amines). In accordance with IUPAC nomenclature, derivatives of ammonium compounds (NH\(_4^+\))\(Y^-\) in which all four of the hydrogens bonded to the nitrogen have been replaced with hydrocarbyl groups, are considered as quaternary ammonium compounds which are not amines. That is, in the amino acids as used in accordance with the present invention the amine group, preferably the amino group, is a RNH\(_2\), R\(_2\)NH or R\(_3\)N residue but not an NR\(_4^+\) residue. Preferably, quaternary ammonium compounds comprising a carboxylic acid group are not used as amino acid additive in accordance with the present invention.

In a preferred embodiment of the invention, the amino acid used as additional additive is a \(\beta\)-amino acid or an \(\alpha\)-amino acid, most preferred an \(\alpha\)-amino acid, \(\alpha\)-amino acids generally have a chemical structure according to formula (1)
or a salt thereof. Residue R may be hydrogen, or an alkyl or an optionally substituted aryl or an optionally substituted heteroaryl group. Preferably, residue R is a C1 - C10 alkyl group, in particular a C1 - C6 alkyl group. Most preferably R is a methyl, propan-2-yl (isopropyl), butan-2-yl, or 2-methyl-propan-1-yl.

In a preferred embodiment, the a-amino acid is selected from the group consisting of positively charged amino acids, such as arginine, histidine, and lysine, negatively charged amino acids such as aspartic acid or glutamic acid, polar uncharged amino acids such as serine, threonine, asparagine or glutamine, or cysteine, selenocysteine, glycine and proline. Particular preferred are amino acids with hydrophobic side chain, such as alanine, valine, isoleucine, leucine, methionine, phenylalanine, tyrosine and tryptophan. Most preferred amino acids used as additives are valine, isoleucine and leucine, leucine being most preferred.

a-amino acids are chiral compounds. Generally, both racemic mixtures of the both enantiomers can be used, as well as compositions enriched in one enantiomer, e.g. the D- or the L-enantiomer. Preferably, racemic mixtures of the amino acids may be used in accordance with one embodiment of the present invention.

Suitable derivatives of amino acids are for example esters, such as esters comprising a hydroxyalkyl residue, in particular a hydroxy C1-4 alkyl residue. Alternatively or additionally the amino acid derivative may be an amide. Suitable salts are for example alkali metal and alkali earth metal salts or salts formed between an acid, such as an inorganic acid or a carboxylic acid with the amino group of the amino acid.

The amino acid may be present in the functionalized particulate bicarbonate according to the present invention in the amount of at least 0.02 % by weight, preferably at least 0.05 % by weight, in particular at least 0.1 % by weight. More than 10 % by weight of the amino acid in the functionalized particulate bicarbonate is disadvantageous for cost reasons. Preferably, at most 8 % by weight, more preferably at most 6 % by weight, in particular at most 5 % by weight of the amino acid are present in the functionalized particulate bicarbonate in accordance with one embodiment of the present invention.
The preferred amino acid used as additional additive in the functionalized particulate bicarbonate is leucine. Leucine may for example be present in the functionalized particulate bicarbonate in an amount of 0.02% by weight to 5% by weight, preferably in an amount of 0.05% by weight to 2% by weight, more preferably in an amount of 0.05% by weight to 0.5% by weight.

In one embodiment, the particulate bicarbonate being additionally functionalized with an amino acid, in particular leucine, is prepared by spray-drying.

In additional or alternate embodiments, the additional additive in the functionalized particulate bicarbonate may comprise or consist of a resin acid, a derivative thereof, or a salt thereof. Generally, the resin acid to be used as additional additive in the functionalized particulate bicarbonate is one of the resin acids as known in the art. The resin acids refer to mixtures of related carboxylic acids, preferably abietic acid, found in tree resins. Typically, resin acids have the basic skeleton of three fused rings with an empirical formula C19H20COOH. Preferred the resin acid is a tricyclic diterpene carboxylic acid, more preferable belonging to the abietane diterpene group. Preferred resin acids are abietic-type acids, e.g. selected from the group consisting of abietic acid (abieta-7, 13-dien-18-oic acid), neoabietic acid, dehydroabietic acid, and palustric acid. Also suitable are pimaric-type acids, selected from the group consisting pimaric acid (pimara-8(14),15-dien-18-oic acid), levopimaric acid, or isopimaric acid. Such acids are available from natural sources or via chemical synthesis as e.g. known from US 2014/0148572 A1.

In the context of the present application, "abietic acid" is also referred to as "rosin acid".

A derivative containing resin acids that may be used in accordance with the present invention is tall oil. Tall oil (also called liquid rosin) is obtained as a by-product of the Kraft process of wood pulp manufacture. Crude tall oil contains rosin, resin acids (mainly abietic acids and its isomers), fatty acids (mainly palmitic, and oleic), fatty alcohols, sterols and alkyl hydrocarbon derivatives. Most preferred abietic acid, dehydroabietic acid, neoabietic acid, pimaric acid and their salts, in particular the sodium salts, respectively are used as additional additive in accordance with the present invention.

The resin acid, derivative thereof, or salt thereof may be present in the functionalized particulate bicarbonate according to the present invention in an amount of at least 0.02% by weight, preferably at least 0.05% by weight, in particular at least 0.1% by weight. The resin acid, derivative thereof or salt thereof, such as rosin acid, may for example be present in an amount from 0.02% by weight to 25% by weight, preferably from 0.02% by weight to 20%
by weight or from 0.1% by weight to 11% by weight, such as from 0.5% by weight to 10% by weight.

The preferred resin acid used as additional additive in the functionalized particulate bicarbonate is resin acid, derivative thereof, or salt thereof.

The rosin acid, derivative thereof, or salt thereof may for example be present in an amount from 1% by weight to 25% by weight, preferably from 5% by weight to 20% by weight in the functionalized bicarbonate.

Suitable derivatives of rosin acid may consist of or comprise for example one or more rosin acid esters, such as Ci-25 alkyl esters of rosin, glycerol rosin esters, pentaerythritol rosin esters, or combinations thereof. Other suitable derivatives of rosin acid may comprise hydrogenated rosin acid (such as dihydroabietic acid), neoabietic acid, dimers of rosin acid, or even polymerized rosin.

A preferred derivative of rosin acid may comprise dihydroabietic acid (CAS No. 1740-19-8) preferably at least 50 wt% of dihydroabietic acid. Resigral 52 is a commercial product which contains at least 52 wt% of dihydroabietic acid; Resigral 52 supplied by LES DERIVES RESINIQUES ET TERPEN IQUES (DRT) in Dax, France is a disproportionated rosin modified by chemical reagents to improve its thermal stability by modifying the proportion of its different isomers.

Other commercial rosin acid-based products manufactured by DRT can be used as one or more sources for the rosin acid derivatives, such as:

- water based rosin dispersions (DERMULSENE RE 1513 = aqueous, solvent free dispersion based on stabilized rosin ester, whose solid content is about 56 wt%);
- DERMULSENE A 75 10 = stabilized and polymerized rosin dispersion,
- glycerol rosin esters (DERTOLINE G2L);
- pentaerythritol rosin esters (DERTOLINE P2L = tall oil resin esterified with pentaerythritol; HYDROGRAL P = pentaerythritol ester of hydrogenated rosin);
- dimeric rosin acid (POLYGRAL 95, POLYGRAL 115, POLYGRAL 140);
- polymerized rosin acid (POLYGRAL); and/or
- hydrogenated rosin (HYDROGRAL).

Liquid rosin esters from DRT may be selected from GRANOLITE TEG = triethylene-glycol rosin ester; GRANOLITE M = methyl ester of rosin; and/or HYDROGRAL M = methyl ester of hydrogenated rosin.

The rosin derivatives may comprise or consist of a fortified rosin, that is, a rosin which has a reacted content of a minor amount but effective amount of an acidic compound
containing the \(-\text{COC}==\text{C}—\) group as fortifying agent such as maleic anhydride, fumaric acid, acetylene dicarboxylic acid and the dehydration products of citric acid, converting at least some of the abietic acid and related compounds to tricarboxylic species.

Suitable salts of rosin acid are for example alkali metal and alkali earth metal salts, but other salts are suitable as well. A preferred salt is the sodium salt.

Rosin acid may also be used as additive in the form of mixtures comprising rosin acid, such as tall oil.

In yet additional or alternate embodiments, the additional additive in the functionalized particulate bicarbonate may comprise or consist of a fatty-acid, derivative thereof (such as esters), or a salt thereof.

Fatty acids as used as additive in the present invention are those fatty acids as known in the art, i.e., a carboxylic acid with an aliphatic residue, which is either a saturated or unsaturated. Preferred, the fatty acid is a compound according to formula (II)

\[
R-\text{COOH}
\]

(II)

wherein R is a saturated or unsaturated C\(_6\)-C\(_{18}\) alkyl group, preferably a C\(_{12}\)-C\(_{18}\) alkyl group, such as a C\(_{12}\)-C\(_{18}\) alkyl group. The fatty acids may be used in form of their salts, in particular sodium or potassium salts, most preferably sodium salt. Even more preferred residue R is a C\(_{16}\)-C\(_{18}\) alkyl group, most preferred, the fatty acid is palmitic acid, linoleic acid, lauric acid or stearic acid, the latter being most preferred.

Examples of fatty acid derivatives are glycerides. Glycerides are esters formed from glycerol and fatty acids, in particular glycerol mono, di and tri fatty esters.

The preferred fatty acid used as additional additive in the functionalized particulate bicarbonate is stearic acid, an ester thereof, or a salt thereof. The more preferred fatty acid used as additional additive in the functionalized particulate bicarbonate is stearic acid, a stearate salt, or an ester of stearic acid, such as its ester with glycerol, such as tristearin, or glyceryl tristearate, which is a triglyceride derived from three units of stearic acid. Another preferred additive is glycerol mono stearate.

The fatty acid, derivative thereof, or salt thereof may be present in the functionalized particulate bicarbonate according to the present invention in an amount of at least 0.02% by weight, preferably at least 0.05% by weight, in particular at least 0.1% by weight. In certain embodiments the fatty acid, derivative thereof, or salt thereof may be present in the
functionalized particulate bicarbonate in an amount of from 0.02% by weight to 30% by weight, preferably from 0.1% by weight to 10% by weight, more preferably from 0.5% by weight to 7% by weight.

In some embodiments, the additive may exclude stearic acid, an ester thereof, or salts thereof.

In yet more additional or alternate embodiments, the additional functionalization additive may comprise or consist of a polymer (other than the polyoxyalkylene additive), such as a polymer selected from the group consisting of polyvinylalcohol, polyglycol, polysaccharide, poly(meth)acrylic acid, poly(acrylic acid co-maleic acid), polyvinylpyrrolidone, N-2(-Hydroxypropyl) methacrylamide, and combinations thereof.

The polymer may be a natural or synthetic polymer. Natural polymers are polymers being from natural sources, such as starch and arabic gum. Natural polymers may also be modified, such as hydrolyzed starch.

Synthetic polymers are for example poly(meth)acrylates and derivatives thereof, and polyvinylalcohols. MetWacrylic polymers may for example be anionic polymers with methacrylic acid as functional groups, cationic polymers with methylaminoethyl methacrylates as functional groups, meth-/acrylate copolymers with trimethyl-aminoethyl-methacrylate as functional groups and neutral polymers of metWacrylates which are available from Evonik under the trade name Eudragit®. Suitable Eudragit® grades are for example grades L, S, FS, E, RL, RS, NE and NM. Preferred are Eudragits® of grade RL, in particular Eudragit® RL 30D.

In some embodiments, when a polysaccharide is used as additional additive in the functionalized particulate bicarbonate, the polysaccharide additive may be at least one polysaccharide selected from the group consisting of hydrolyzed starch, carboxymethyleellulose, alginic acid and salts thereof, arabic gum, carrageenan, guar gum, locust bean gum, xantham gum, and combinations thereof.

The additional functionalization additive in the functionalized particulate bicarbonate may comprise or consist of a polysaccharide selected from the group consisting of:

- guar gums and their derivatives, in particular hydroxypropyl guar (such as Jaguar HP-105);
- alginic acid and its salts, such sodium, calcium or copper (e.g., Kaltostat, Calginat, Landalgine. Kalrostat, Kelacid, Vocoloid, Xantalgin); and
carboxymethylcelullose (e.g., Aquaplast, Carmethose, CELLOFAS, Cellpro, Cellugel, Collowel, Ethoxose, Orabase, Lovosa). 

In another embodiment, the additional functionalization additive of the functionalized particulate bicarbonate may comprise or consist of a modified, in particular hydrolyzed starch or a compound comprising such starch. Particularly preferred additives of this class are hydrolyzed starch, arabic gum and maltodextrin, maltodextrin being particularly preferred.

The additional functionalization additive may be present in the functionalized particulate bicarbonate according to the present invention in an amount of at least 0.02% by weight, preferably at least 0.05% by weight, more preferably at least 0.1% by weight. In particular the hydrolyzed starch, arabic gum, maltodextrin, poly(meth)acrylate, and polyvinylalcohol may for example be present in an amount of between 0.02% by weight to 40% by weight, more preferably from 0.1% by weight to 35% by weight, even more preferably from 1% by weight to 20% by weight, such as from 2% by weight to 10% by weight.

In one embodiment the functionalized particulate bicarbonate comprises at least one polymer as additional functionalization additive together with the polyoxyalkylene additive, wherein the polymer is different than the polyoxyalkylene additive. In this combination the polymer preferably is selected from the group consisting of poly(meth)acrylates and derivatives thereof, polyvinyl alcohol and polysaccharides, including modified, in particular hydrolyzed starch, maltodextrin and arabic gum.

In a specific embodiment of a combination of the polyoxyalkylene additive (such as polyethylene glycol, preferably low molecular weight (<1000) polyethylene glycol) and rosin acid or any derivative or salt thereof as functionalization additives of the functionalized particulate bicarbonate, the rosin acid is present in an amount of from 5% by weight to 20% by weight, preferably of from 1% by weight to 11% by weight of the total amount of functionalized particulate bicarbonate, and the polyoxyalkylene additive (such as polyethylene glycol) is present in an amount of from 1% by weight to 30% by weight, preferably of from 5% by weight to 25% by weight, such as for example from about 10% by weight to about 20% by weight, each of the total amount of the functionalized particulate bicarbonate.

In a further preferred embodiment, the functionalized particulate bicarbonate comprising the polyoxyalkylene additive (in particular, polyethylene glycol) as one functionalization additive and rosin acid or any derivative or salt thereof as another
functionalization additive is prepared by first co-milling bicarbonate particles in the presence of the rosin acid or derivative or salt thereof, and the thus obtained functionalized particles are then mixed with the polyoxyalkylene additive (such as polyethylene glycol) and extruded to obtain particulate bicarbonate being functionalized with rosin acid and the polyoxyalkylene additive, in particular polyethylene glycol. The resulting extruded functionalized particulate bicarbonate may be further subjected to milling to reduce the mean particle size of the final product.

Oils used as additional additive in the present invention may be organic oils or mineral oils which may be of animal, vegetable, or petrochemical origin. Suitable oils are for example olive oil, maize oil, sunflower oil and soybean oil.

The oil may be chemically modified, such as epoxidized. A preferred oil is epoxidized soybean oil. The oil may be present in the functionalized particulate bicarbonate according to the present invention in an amount of at least 0.02% by weight, preferably at least 0.05% by weight, in particular at least 0.1% by weight. In preferred embodiments the oil can be present in the functionalized particulate bicarbonate in an amount of from 0.1% by weight to 10% by weight, more preferably from 1% by weight to 7% by weight.

In a further embodiment the additional additive in the functionalized particulate bicarbonate is a wax, such as bees wax or carnauba wax.

The wax may be present in the functionalized particulate bicarbonate according to the present invention in the amount of at least 0.02% by weight, preferably at least 0.05% by weight, in particular at least 0.1% by weight. The wax can for example be present in an amount of from 1% by weight to 30% by weight, preferably from 5% by weight to 25% by weight.

Additional information on particular properties of functionalized particulate bicarbonate can be found in WO2016/102591A1 by SOLVAY SA when the additive comprises or consists of an amino acid or salt thereof; in EP3037388A1 by SOLVAY SA when the additive comprises or consists of a resin acid or fatty acid as additive.

In some embodiments, the functionalized bicarbonate may further comprise a processing aid, an anti-caking agent and/or a flow aid.

This processing aid, anti-caking agent and/or flow aid may be first mixed with the functionalization additive before their mixture is used to functionalize the particulate bicarbonate. Alternatively, this processing aid, anti-caking agent and/or flow aid may be mixed with the functionalized bicarbonate. For example, the functionalized bicarbonate powder may be mixed in a Lodge-type “ploughshare” mixer by introducing the processing
aid, anti-caking agent and/or flow aid progressively.

In some preferred embodiments, the functionalized bicarbonate may further comprise silica. Silica may be used as a processing aid, an anti-caking agent and/or a flow aid for the functionalized bicarbonate. It is recommended that the silica is in amorphous (and noncrystalline) form. Preferably the silica in the functionalized bicarbonate is amorphous precipitated silica. The functionalized bicarbonate may comprise at least 0.1 wt%, preferably at least 0.2 wt%, preferably at least 0.5 wt% of silica based on the total composition of the functionalized bicarbonate. It is recommended that the functionalized bicarbonate comprises not more than 5 wt%, preferably not more than 4 wt% silica, more preferably not more than 3 wt% silica. The presence of silica can improve the flow of the functionalized bicarbonate and therefore promotes its homogeneous application as a chemical foaming agent in a foamy polymer composition.

It is preferable that the silica is in the form of very fine particles, having a high specific surface area, for example greater than 180 m²/g, preferably at least 200 m²/g, more preferably at least 400 m²/g, measured according to the ISO 5794-1 standard, annex D.

According to an advantageous embodiment, the silica in the functionalized bicarbonate is in the form of particles having a mean diameter of less than 20 µm, preferably having a mean diameter of at most 15 µm, or at most 10 µm, more preferably at most 8 µm. The mean diameter is measured according to the ASTM C-690-1992 standard.

Suitable commercially available silicas are: amorphous silica Sipernat® 50S, manufactured by the company Degussa; amorphous silica Sipernat 22 S produced by Degussa, having a lower specific surface of 19(m²/g instead of 450 m²/g for Sipernat 50S; silica Tixosil® 38AB produced by Rhodia (owned by Solvay) having a larger mean diameter of 15µm (instead of 7µm for Sipernat 50S and 22S). Tixosil® anti-caking agents from Rhodia with their controlled particle size are effective flow-conditioning agents for powders adversely affected by moisture or humidity, pressure, or static charge.

In such embodiments, the functionalized bicarbonate may comprise from 0.1 wt% to 5 wt% amorphous silica, preferably from 0.2 wt% to 4 wt% amorphous silica, more preferably from 0.5 wt% to 3 wt% amorphous silica. The silica is preferably precipitated silica.

In such embodiments, the functionalized bicarbonate may comprise a weight ratio of sodium bicarbonate to silica of from 12:1 to 990:1, preferably from 14:1 to 99:1, more preferably from 16:1 to 97:1.

METHOD OF MAKING FUNCTIONALIZED PARTICULATE BICARBONATE
The present invention further relates to a method for making functionalized particulate bicarbonate.

The functionalized particulate bicarbonate may be prepared from a solution containing a bicarbonate ingredient or directly from an already-formed particulate bicarbonate.

The bicarbonate ingredient or particulate bicarbonate may comprise or consists essentially of ammonium bicarbonate, sodium bicarbonate, potassium bicarbonate, or mixtures thereof, in particular the bicarbonate ingredient or particulate bicarbonate comprises or consists essentially of sodium bicarbonate. The bicarbonate ingredient or particulate bicarbonate preferably comprises at least 80 wt% of sodium bicarbonate.

The functionalized particulate bicarbonate according to one aspect of the present invention may be obtained by encapsulation or coating processes.

In particular, another aspect of the present invention relates to a process for preparing functionalized particulate bicarbonate, preferably a functionalized particulate alkali metal bicarbonate as described above according to the invention by at least one of the following processes:

- by spray-drying (also known as atomization), wherein the additive is dissolved in the bicarbonate-containing solution.
- by grinding or co-grinding (also known as milling or co-milling) with the additive(s) in emulsion or powder form;
- by spray coating and granulation within a fluidized bed,
- by spray agglomeration within a fluidized bed,
- by spray chilling (e.g., spray cooling, spray freezing),
- by roller compaction, and/or
- by extrusion, including simultaneous mixing/extrusion.

Spray-drying or drying by atomization is a drying technique. This method comprises spraying the product to be dried, which is in the form of a solution (or a suspension) in a stream of hot gas, so as to obtain a powder in a few seconds or fractions of seconds. The separation of a solution into fine droplets gives rise to a large material transfer surface and it leads to rapid evaporation of the solvent of the solution used.

Suitable apparatus for spray-drying are known in the art, and generally comprise several modules: a module comprising a circuit for storing and atomizing the solution comprising equipment for atomizing or spraying the solution, a module for the preparation of hot gas and its transfer to a drying chamber where it comes into contact with the sprayed
solution, a drying chamber where the sprayed solution is evaporated and the particles are formed, and a module for collecting the particles, generally comprising a cyclone and/or a suitable filter.

Generally, the equipment for atomizing or spraying the solution is a compressed gas sprayer or a dispersion turbine. Also ultrasound nozzles can be used for spraying the solution.

In the spray-drying process of the invention generally an aqueous solution of the bicarbonate is used. While other polar solvents or mixtures of polar solvents, for examples mixtures of water and ethanol, in which the additive is soluble, may be used, water is the preferred solvent.

In the spray-drying method of the present invention the aqueous solution to be spray-dried comprises 1 to 10 % by weight of the bicarbonate ingredient. The bicarbonate ingredient in the solution is preferably an alkali metal bicarbonate. The solution to be spray-dried further comprises 1 to 10,000 ppm of an additive or salt thereof. The additive used is preferably one of those as described above for the particulate bicarbonate of the present invention. In preferred embodiments, the content of the additive in the solution to be spray-dried is 1 to 5,000 ppm, more preferred 1 to 3,000 ppm, in particular 10 to 2,000 ppm, e.g. 50 - 1,000 ppm of additive per kg of solution to be spray-dried. Generally, the aqueous solution comprises at least 1 mg, preferably at least 5 mg, more preferably at least 10 mg, even more preferred at least 100 mg of the additive per kg of aqueous solution. Generally, the aqueous solution comprises at most 2,000 mg, preferably at most 1,500 mg, more preferably at most 1,200 mg of the additive per kg of aqueous solution. In case of salts, weight percentages are given based on the free base / acid.

Generally, in the spray-drying process in the method for making functionalized particulate bicarbonate, the aqueous solution comprises at least or more than 1 %, preferably at least or more than 2 %, more preferably at least or more than 3 %; even more preferably at least or more than 4 %, in particular at least or more than 5 % by weight of the bicarbonate ingredient. Preferably, the bicarbonate ingredient comprises or consists essentially of sodium bicarbonate and/or potassium bicarbonate, in particular comprise or consists essentially of sodium bicarbonate. A high concentration of the bicarbonate ingredient in the aqueous solution is detrimental as leading to high speed plugging of the spraying or atomizing device. Therefore it is generally recommended that the aqueous solution comprises at most or less than 10 % by weight, preferably at most or less than 8 % by weight, more preferably at most or less than 6 % by weight of the bicarbonate ingredient, in particular sodium bicarbonate. Preferably, the bicarbonate-containing solution is an aqueous solution comprising 1 % - 10
%, advantageously 3% - 8%, more advantageously 4% - 8% by weight, such as 4% - 6% by weight of the bicarbonate ingredient, in particular sodium bicarbonate.

The drying with a hot gas breaks part of the alkali metal bicarbonate down into the form of sodium carbonate, CO₂ and water. In one advantageous embodiment of the present invention, the spray-drying is carried out in a gas comprising at least 5%, advantageously at least 10%, more advantageously at least 20%, and even more advantageously at least 30% of CO₂ by volume on a dry gas base. This enables to limit the bicarbonate decomposition into carbonate solid and CO₂ gas and water vapor. Generally, the spray-drying is carried out with a gas preheated between 40°C and 220°C. Advantageously the spray-drying is carried out in a spray-drying chamber and wherein the gas is preheated before being introduced into the spray-drying chamber at least 40°C, preferably at least 50°C, more preferably at least 60°C, even more preferably at least 70°C. Also advantageously, the gas is preheated before being introduced into the spray-drying chamber at most 220°C, preferably at most 200°C, more preferably at most 180°C, even more preferably at most 130°C.

It is preferable for the temperature of the gas after the spray-drying operation to be at most 80°C, advantageously at most 70°C and more advantageously at most 60°C.

In one embodiment in the method for making functionalized particulate bicarbonate, the aqueous solution is preheated to a temperature of at least 20°C and preferably at most 80°C before being sprayed during the spray-drying operation. In one particular embodiment, the aqueous solution is preheated to a temperature of at least 20°C and at most 25°C before being sprayed during the spray-drying operation.

The method for making functionalized particulate bicarbonate according to the present invention may comprise co-grinding of the bicarbonate ingredient in the presence of the additive, such as in the presence of 0.02 to 10 parts by weight of an additive per 100 parts per weight of the substance undergoing co-grinding. The bicarbonate ingredient and the additive are preferably as defined above.

In the process for preparing the functionalized particulate bicarbonate by co-grinding, all suitable grinding procedures as known in the art can be used.

Typical devices include impact mills, which are mills in which the material be milled as subjected to the impact of moving mechanical part and that have the effect of fragmenting the particles of the material. Impact mills are well-known in the fine milling art. Such mills include hammer mills, spindle mills, attritor mills, jet mills, ball mills, such as planetary ball mills, and cage mills. Such mills are e.g. manufactured and available by Grinding Technologies and System SRL or by Hosokawa Alpine AG. Most preferred, an Alpine LGM
3 is used. In the process for preparing the alkali metal bicarbonate particles, the alkali metal bicarbonate is grinded in the presence of the additive, i.e. the amino acid as defined above. Either the total amount of bicarbonate and additive is added into the mill at once, followed by milling or preferably the bicarbonate and the additive are fed into the milling device at a constant rate. Suitable rates for the bicarbonate are 50 kg/h to 500 kg/h, preferably 100 kg/h to 400 kg/h, e.g. about 150 kg/h. The amount of additive corresponds to the weight ratio of the bicarbonate ingredient and additive used. For examples, if the bicarbonate ingredient is co-grinded in the presence of 1 part by weight of an additive, per 100 parts by weight of the substance undergoing co-grinding, the feeding rate of the additive is only 1% of the feeding rate of the bicarbonate ingredient.

The amount of functionalization additive (e.g., polyoxyalkylene additive, additional additive as previously described) in the process for preparing the functionalized particulate bicarbonate by co-grinding, is from 0.02 to 10 parts by weight per 100 parts by weight of the substance undergoing co-grinding. Below 0.02 parts by weight, there is only low efficacy of the additive. Using higher amounts than 10 parts by weight of the additive is disadvantageous for cost reasons. Preferred amounts are 0.2 - 8 parts by weight of the additive, more preferred 0.5 - 5 parts by weight of additive, even more preferred 0.8 - 2 parts by weight of additive, in particular about 1 part by weight of additive, each per 100 parts by weight of the substance (typically the bicarbonate ingredient and the at least one additive) undergoing co-grinding.

In some embodiment, the method for making functionalized particulate bicarbonate according to the present invention may comprise spray coating, for example within a fluidized bed. Spray coating within a fluidized bed is a technique in which a powder (solid particles of the bicarbonate ingredient) is fed into a fluidization chamber. A gas fluidizes the powder from the chamber bottom through a grid. A liquid which comprises the functionalization additive in dissolved form, in a melted form, and/or in dispersed solid form (for example a solution, an emulsion, a suspension, a melt, a melt emulsion or a melt suspension) is sprayed into the fluidized powder to apply a layer or coating onto particles.

In some embodiment, the method for making functionalized particulate bicarbonate according to the present invention may comprise spray granulation, for example within a fluidized bed. The fluidized bed spray granulation is a method for making free-flowing granulate from liquids. The liquid containing solids, such as aqueous solutions, emulsions, suspensions, melts, melts emulsions or melts suspensions, is sprayed into a fluidized bed system. The solids are preferably bicarbonate particles. Due to the high heat exchange the aqueous or organic solvent in the liquid evaporates immediately, and the solids form small
particles as starter cores. These are sprayed with another liquid with contains the additive (solution/suspension). A gas fluidizes the additive solution/suspension sprayed into a chamber. After evaporation and drying in the fluidized bed, the dried additive forms a hard coating around the starter core. This step is continuously repeated in the fluidized bed so that the granulate grows to form onion-like or blackberry-like structures. An onion-like structure is obtained from layer by layer coating. Alternatively, a defined volume of suitable starter cores can be provided. In this option, the liquid only serves as a vehicle for the solids that are being applied.

In some embodiment, the method for making functionalized particulate bicarbonate according to the present invention may comprise spray agglomeration, for example within a fluidized bed. Spray agglomeration within a fluidized bed is a technique in which a powder or a fine granulate is fed into a fluidization chamber. A gas fluidizes the powder from the chamber bottom through a grid. A liquid (either a solution, an emulsion, a suspension a melt, a melt emulsion or a melt suspension), which acts as a binder, is sprayed on the fluidized particles. Liquid bridges are created that form agglomerates from the particles. Spraying continues until the desired size of the agglomerates is reached.

In some embodiment, the method for making functionalized particulate bicarbonate according to the present invention may comprise spray chilling (or spray cooling, spray freezing). Spray chilling is a technique in which a melt, a melt emulsion or a melt suspension is sprayed into a fluidization chamber. A cold gas in injected in the fluidization chamber. The solidification of solid particles is achieved by the melt droplets losing heat to cold air in the fluid bed.

In some embodiment, the method for making functionalized particulate bicarbonate according to the present invention may comprise roller compaction. Roller compaction is a technique in which powder particles are made to adhere to each other by applying a force onto the powder, which causes a considerable size enlargement. The powder is compacted between two counter rotating rolls to apply a force. The obtained briquettes, flakes or ribbons are crushed out of the rolls to reach a desired particle size.

In some embodiment, the method for making functionalized particulate bicarbonate according to the present invention may comprise extrusion (or mixing extrusion). Extrusion (or mixing extrusion) is a technique in which a powder or another material is pushed through a die of a fixed cross-section. A screw, two screws or a succession of paddles may help pushing the material through mixing, degassing and homogenizing phases. Temperature
control along the distance allows for phase changing, melting, crystallization, chemical reaction, coating or granulating the materials.

In some embodiments when the functionalized bicarbonate further comprises silica, preferably amorphous silica, more preferably amorphous precipitated silica, silica may be first mixed with the functionalization additive before their mixture is used to functionalize the particulate bicarbonate. Alternatively, silica may be mixed with the functionalized bicarbonate after it is prepared. For example, the functionalized bicarbonate powder may be mixed in a Lodige-type "ploughshare" mixer by introducing the silica progressively into the functionalized bicarbonate. Silica may be used in the functionalized bicarbonate as a processing aid, an anti-caking agent (such as acting as a dessicant for water capture), and/or a flow aid. For example the functionalized bicarbonate powder may be mixed with amorphous precipitated silica to form a free-flowing powder.

**USE OF THE FUNCTIONALIZED PARTICULATE BICARBONATE**

The present invention further relates to the use of the herein described functionalized particulate bicarbonate as chemical blowing agent for foaming

- a thermoplastic polymer, for example PVC plastisol; or
- a polymer resin in an extrusion process.

**CHEMICAL BLOWING AGENT**

The present invention further relates to a chemical blowing agent for foaming

- a thermoplastic polymer; or
- a polymer resin in an extrusion process.

Thus, another aspect of the invention provides a chemical blowing agent for foaming a thermoplastic polymer which comprises the functionalized particulate bicarbonate as described in the various embodiments herein, wherein the functionalized particulate bicarbonate contains at least one additive as described in the various embodiments herein. Optionally, the functionalized particulate bicarbonate may further comprise a processing aid, an anti-caking agent (such as acting as a dessicant for water capture), and/or a flow aid. In some embodiments, the functionalized particulate bicarbonate may further comprise silica, preferably amorphous silica, more preferably amorphous precipitated silica, as described in the various embodiments herein. In some embodiments, the chemical blowing agent for
foaming a thermoplastic polymer comprises a functionalized particulate bicarbonate having a particle size \( D_{50} \) of 1000 nm or less.

In alternate embodiments, the chemical blowing agent for foaming a thermoplastic polymer comprises a functionalized particulate bicarbonate having a particle size \( D_{50} \) more than 1 micron and up to 250 \( \mu \)m, preferably up to 100 \( \mu \)m, more preferably up to 60 \( \mu \)m, yet more preferably up to 40 \( \mu \)m, or up to 30 \( \mu \)m, or up to 25 \( \mu \)m.

The additive in the functionalized particulate bicarbonate is as defined in the various embodiments described above.

Any particular embodiments for the functionalized particulate bicarbonate and the additive(s) used for its functionalization described herein are applicable here.

In some embodiments, the chemical blowing agent does not contain a compound which liberates nitrogen gas during heating.

In some embodiments, the chemical blowing agent does not contain a compound which liberates ammonia during heating.

In preferred embodiments, the chemical blowing agent is an endothermic chemical blowing agent comprising the functionalized particulate bicarbonate as described in the various embodiments herein.

In preferred embodiments, the chemical blowing agent does not contain an exothermic blowing agent.

In some embodiments, the chemical blowing agent comprises the functionalized particulate bicarbonate and further comprises a second compound as another blowing agent.

The second compound is preferably an endothermic blowing agent.

The second compound preferably can liberate CO\(_2\) upon heating. This second compound preferably increases the CO\(_2\) generation which is already formed by the decomposition of the functionalized particulate bicarbonate upon heating.

This CO\(_2\)-liberating second compound acting as blowing agent may be a carboxylic or polycarboxylic acid, derivative thereof (such as esters), or salts thereof.

Suitable carboxylic acids include those of the formula: \( \text{HOOC—R—COOH} \) where R is an alkylene group of 1 to about 8 carbon atoms which may also be substituted by one or more hydroxy groups or keto groups and may also contain unsaturation. Also included are esters, salts and half salts.

A preferred CO\(_2\)-liberating second compound acting as blowing agent may include at least one of:

- fumaric acid,
- tartaric acid, or
- citric acid, citrates (such as sodium hydrogen citrate, disodium citrate), or esters of citric acid.

Esters of citric acid may include tributyl citrate, triethyl citrate, tri-C12-13 alkyl citrate,
tri-C14-15 alkyl citrate, tricaprylyl citrate, triethylhexyl citrate, triisocetyl citrate,
triocytldodecyl citrate and trisostearyl citrate, isodecyl citrate and stearyl citrate, dilauryl citrate,
and/or ethyl citrates (mixture of tri-, di- and monoesters), preferably tributyl citrate,
triethyl citrate, isodecyl citrate, or triethylhexyl citrate.

A more preferred C02-liberating second compound acting as endothermic blowing agent comprises or consists of citric acid, esters thereof, or salts thereof.

In yet alternate embodiments, the chemical blowing agent may further comprise an exothermic chemical blowing agent, preferably selected from the group consisting of ADCA (azodicarbonamide), OBASH (4,4’-oxybis(benzenesulfonylhydrazide)), DNPT (dinitrosopentamethylene tetramine), PTSS (p-toluene semicarbazide), BSH (benzene-4-sulfonohydrazide), and TSH (toluene-4-sulfonohydrazide), 5-PT (5-phenyltetrazole), more preferably may further comprise ADCA (azodicarbonamide). In such embodiments, the chemical blowing agent may comprise a weight ratio of the functionalized bicarbonate to the exothermic chemical blowing agent from 5:95 to 95:5, preferably 90:10 to 10:90, more preferably 85:15 to 15:85, yet more preferably 80:20 to 20:80.

Optionally, the chemical blowing agent comprising the functionalized particulate bicarbonate may further comprise a processing aid, an anti-caking agent (such as acting as a dessicant for water capture), and/or a flow aid. In some embodiments, the chemical blowing agent comprises a mixture of the particulate functionalized particulate bicarbonate and silica, preferably amorphous silica, more preferably amorphous precipitated silica.

FUNCTIONALIZED SECOND COMPOUND IN BLOWING AGENT COMPOSITION

It is also envisioned that the C02-liberating second compound in the blowing agent composition which supplements C02 generation may also be in a functionalized particulate form. This functionalized particulate second compound would comprise an additive as described herein in relation to the functionalized particulate bicarbonate. This functionalization of the particulate second compound may also use one or more techniques as described above in relation to the making of the functionalized particulate bicarbonate.
In some embodiments, the endothermic blowing agent comprises, or consists essentially of, or consists of:
- the functionalized particulate bicarbonate, as described herein in the various embodiments; and
- a functionalized particulate carboxylic or polycarboxylic acid, derivative thereof (such as esters), or salts thereof.

In particular embodiments, the endothermic blowing agent comprises, or consists essentially of, or consists of:
- the functionalized particulate bicarbonate, as described herein in the various embodiments; and
- a functionalized particulate carboxylic or polycarboxylic acid, derivative, or salt thereof, selected from the group consisting of: fumaric acid, tartaric acid, citric acid, salts thereof (such as sodium hydrogen citrate, disodium citrate), esters thereof, and any combination thereof.

In some embodiments, the chemical blowing agent for foaming a thermoplastic polymer comprises a functionalized particulate bicarbonate and a functionalized particulate second compound (e.g., functionalized carboxylic or polycarboxylic acid, derivative, or salt thereof), both having a particle size D50 of 1000 nm or less.

In alternate embodiments, the chemical blowing agent for foaming a thermoplastic polymer comprises a functionalized particulate bicarbonate and a functionalized particulate second compound (e.g., functionalized carboxylic or polycarboxylic acid, derivative, or salt thereof), both having a particle size D50 more than 1 micron and up to 250 microns, preferably up to 100 microns, more preferably up to 60 microns, still more preferably up to 40 microns or up to 30 microns.

In yet alternate embodiments, the chemical blowing agent for foaming a thermoplastic polymer comprises a functionalized particulate bicarbonate having a particle size D50 of 1000 nm or less and a functionalized particulate second compound (e.g., functionalized carboxylic or polycarboxylic acid, derivative, or salt thereof) having a particle size D50 more than 1 micron and up to 250 microns, preferably up to 100 microns, more preferably up to 60 microns, still more preferably up to 40 microns or up to 30 microns.

In embodiments where the endothermic blowing agent comprises the functionalized particulate bicarbonate and the functionalized particulate second compound (e.g.,
functionalized carboxylic or polycarboxylic acid, derivative, or salt thereof), the method for making such blowing agent may contain the following steps:

- **A/** making the functionalized particulate bicarbonate using a first additive (wherein the first additive is selected from those defined herein); separately making the functionalized particulate second compound using a second additive (wherein the second additive is selected from those defined herein); mixing the functionalized particulate bicarbonate and the functionalized particulate second compound to make the endothermic blowing agent (so long as one of the first and second additives is the polyoxyalkylene additive); or

- **B/** mixing the particulate bicarbonate and the particulate second compound before functionalization to make a non-functionalized particulate mixture; and functionalizing the non-functionalized particulate mixture using at least one additive (comprising at least the polyoxyalkylene additive and optionally one or more additional functionalization additives as defined herein) to make the endothermic blowing agent.

In the method **A/**, the first and second additives may be the same or may be different.
The techniques to make the functionalized particulate bicarbonate and the functionalized particulate second compound may be the same or may be different.

In the method **B/**, the additive and the technique for functionalization of the particulate bicarbonate and the particulate second compound is generally the same, thus resulting in less flexibility in the making of the endothermic blowing agent. However this method would be more cost effective.

**FOAMABLE POLYMER COMPOSITION**

Another aspect of the present invention relates to a foamable polymer composition comprising

a) a polymer;

b) the functionalized particulate bicarbonate used as chemical blowing agent for foaming; and

c) at least one polymer additive.

Optionally, the functionalized particulate bicarbonate in the foamable polymer composition may further comprise a processing aid, an anti-caking agent (such as acting as a dessicant for water capture), and/or a flow aid. In some embodiments, the functionalized particulate bicarbonate in the foamable polymer composition may further comprise silica.
preferably amorphous silica, more preferably amorphous precipitated silica, as described in
the various embodiments herein.

The foamable polymer composition may optionally comprise a functionalized
particulate second compound used as supplemental chemical blowing agent for foaming (as
described herein).

The foamable polymer composition may optionally comprise a foam stabilizer, such as
a silicone based foam stabilizer.

The amount of the polymer in the foamable polymer composition typically ranges
between about 10 wt % (or 20 wt%) and about 90 wt % based on the total weight of the
foamable polymer composition. The amount of polymer can be selected according to the
desired final characteristic of the foamable polymer composition.

In one preferred embodiment according to the present invention, the foamable polymer
composition is in the form of a plastisol. The term "plastisol" relates to a suspension of
polymer particles in a liquid plasticiser.

Preferably, the plastisol is a suspension of polyvinyl chloride particles in a liquid
plasticizer. The choice of the plasticizer is not particularly limited and, therefore, commonly
employed plasticizers, such as dioctylphthalate or 1,2-cyclohexyldicarbonic acid diisononyl
ester can be used for this purpose.

The terms "polyvinyl chloride" intend to designate vinyl chloride homopolymers as
well as copolymers of vinyl chloride with other ethylenically unsaturated monomers which
are either halogenated (chloroolefins like vinylidene chloride; chloroacrylates; chlorinated
vinylethers) or non halogenated (olefins like ethylene and propylene; styrene; vinylethers like
vinyl acetate) monomers; as well as vinyl chloride copolymers with acrylic and methacrylic
acids; esters, nitriles and amides. Vinyl chloride homopolymers and vinyl chloride
copolymers containing 50-99 weight %, preferably 60-85 weight % of vinyl chloride are
preferred.

The amount of the plastisol in the foamable polymer composition typically ranges
between about 20 wt % and about 90 wt % based on the total weight of the foamable polymer
composition. The amount of plastisol can be selected according to the desired final
characteristic of the foamable polymer composition.

The amount of the functionalized particulate bicarbonate in the foamable polymer
composition typically ranges from 5 to 15 parts per 100 parts by weight of polymer. For
example, for a foamable PVC polymer composition, the amount of the functionalized
particulate bicarbonate in the foamable PVC polymer composition typically ranges from 5 to
15 parts per 100 parts by weight of PVC resin, preferably from 6 to 12 parts per 100 parts by weight of PVC resin; more preferably from 7 to 11 parts per 100 parts by weight of PVC resin or from 8 to 10 parts per 100 parts by weight of PVC resin.

The foamable polymer composition of the present invention may contain further components, for instance, foam stabilizers; air release agents; fillers or extenders (e.g., CaCO₃ as a filler), such as carbon black; other polymers and oils; curing agents, such as sulfur compounds and various chemicals that act as a part of a curing system, such as zinc oxide; antistatic agents; biocides; colorants; coupling agents; fibrous reinforcements; flame retardants; fungicides; heat stabilizers; lubricants; mold release agents; plasticizers (e.g., DINP = di-isononyl phtalate); preservatives; processing aids; slip agents; ultraviolet stabilizers; viscosity depressants; and any other ingredient that may be a desirable component of the resulting foam polymer.

A preferred foam stabilizer may include a silicone based foam stabilizer, such as BYK 8020 for PVC plastisol.

In some embodiments, the foamable composition does not contain another blowing agent other than the functionalized particulate bicarbonate.

In other embodiments, the foamable composition does not contain any blowing agent which is exothermic. In particular embodiments, the foamable composition does not contain a blowing agent which liberates nitrogen gas and/or ammonia. Examples of blowing agents which liberate nitrogen gas are exothermic blowing agents such as azodicarbonic acid diamide (azodicarbonamide, ADC, ADCA, CAS No. 123-77-3), the sulfonhydrazides 4,4’-oxybis(benzenesulfonylhydrazide) (OBSH, CAS No. 80-51-3) and p-toluensulfonylhydrazide, (TSH, CAS No. 1576-35-8).

In preferred embodiments, the foamable polymer composition does not contain azodicarbonamide.

In alternate or additional preferred embodiments, the foamable polymer composition does not contain benzenesulfonylhydrazide.

In alternate or additional embodiments, the foamable polymer composition does not contain p-toluensulfonylhydrazide.

In yet alternate embodiments, the foamable polymer composition may further comprise an exothermic chemical blowing agent, preferably selected from the group consisting of ADCA (azodicarbonamide), OBSH (4,4’-oxybis(benzenesulfonyl hydrazide)), DNPT (dinitrosopentamethylene tetramine), PTSS (p-toluene semicarbazide), BSH (benzene-4-sulfonohydrazide), and TSH (toluene-4-sulfonohydrazide), 5-PT (5-phenyltetrazole), more
preferably may further comprise ADCA (azodicarbonamide). In such embodiments, the foamable polymer composition may comprise a weight ratio of the functionalized bicarbonate to the exothermic chemical blowing agent from 5:95 to 95:5, preferably 90:10 to 10:90, more preferably 85:15 to 15:85, yet more preferably 80:20 to 20:80.

**PROCESS FOR PREPARING THE FOAMABLE COMPOSITION**

Another aspect of the present invention relates to a process for preparing the foamable polymer composition as described above, wherein the process comprises:

- mixing the components of the foamable polymer composition as provided herein,

wherein a component of the foamable polymer composition is the functionalized particulate bicarbonate.

The process for preparing the foamable polymer composition can be carried out in conventional equipment, for example made of Austenitic Stainless Steel (304L-316L etc). The process can also be carried out in non-metallic materials like plastics, glass and ceramics for chemical use.

Preferably, the mixing step in the process is carried out under continuous mixing. Because the foamable polymer composition has a relatively high viscosity, a powerful mixing is required for obtaining a homogeneous foamable polymer composition. Moreover, because the blowing effect already starts taking place during the mixing, the speed of mixing should be sufficiently high in order to achieve a good homogenisation within a short period of time. Preferably, the homogenisation is achieved within less than about 20 seconds during the mixing step(s) of the process. The selection of the mixing conditions depends upon the specific characteristics of the equipment. The mixing step(s) of the process for preparing the foamable polymer composition can be carried out at any suitable temperature as in conventional foam production, the process can for example be carried out at relatively low temperatures (ambient) ranging between about 0°C and about 30°C, for instance at about 20-25°C.

A foamable polymer composition may be prepared by mixing

- the functionalized particulate bicarbonate in powder form as described herein,
- optionally, an exothermic blowing agent, such as ADCA (azodicarbonamide), OBSD (4,4'-oxybis(benzenesulfonyl hydrazide)), DNPT (dinitrosopentamethyl ene tetramine), PTSS (p-toluene semicarbazide), BSH (benzene-4-sulfonohydrazide), and TSH (toluene-4-sulfonohydrazide), 5-PT (5-phenyltetrazole), and optionally a kicker for the exothermic blowing agent;
at least one plasticizer,
- at least one foamable polymer,
- optionally a foam stabilizer; and
- optionally one or more polymer additives such as an inorganic filler.

For example, a PVC plastisol composition may be prepared by mixing
- the functionalized particulate bicarbonate in powder form as described herein,
- optionally, an exothermic blowing agent, such as ADCA (azodicarbonamide), OBSH (4,4'-oxybis(benzenesulfonyl hydrazide)), DNPT (dinitrosopentamethylene tetramine), PTSS (p-toluene semicarbazide), BSH (benzene-4-sulfonylhydrazide), and TSH (toluene-4-sulfonohydrazide), 5-PT (5-phenyltetrazole) and optionally a kicker for the exothermic blowing agent;
- at least one plasticizer,
- at least one polyvinyl chloride resin,
- optionally a foam stabilizer; and
- optionally one or more polymer additives such as an inorganic filler.

A preferred foam stabilizer may include a silicone based foam stabilizer, such as BYK 8020 for PVC plastisol.

The components of the foamable polymer composition such as a PVC foamable plastisol composition may be dispersed on a high speed mixer and then de-aerated under vacuum.

The PVC foamable plastisol composition may optionally comprise a functionalized particulate second compound used as supplemental chemical blowing agent for foaming (as described herein). The functionalized particulate second compound may be a functionalized particulate carboxylic or polycarboxylic acid, derivative, or salt thereof. The functionalized particulate second compound may be selected from the group consisting of: fumaric acid, tartaric acid, citric acid, salts thereof (such as sodium hydrogen citrate, disodium citrate), esters thereof, and any combination thereof.

**PROCESS FOR PREPARING A FOAMED POLYMER**

A further aspect of the present invention relates to a process for preparing a foamed polymer, wherein the above described foamable polymer composition is heated. Thus, the process for preparing a foamed polymer can comprise the following step:
- mixing the components of the foamable polymer composition as provided herein; and
- heating the foamable polymer composition obtained from mixing.
Preferably, the heating step is carried out immediately after the completion of the mixing step.

The process for preparing a foamed polymer may include a shaping step such as extrusion or spreading. Preferably, the heating step is carried out immediately after the completion of the mixing and shaping steps.

The heating step can be carried out at a temperature from about 120°C and up to about 220°C, or from about 120°C and up to about 210°C, or from about 120°C and up to about 200°C, depending on the resin and the composition of plastisol.

The time of heating in the heating step depends on the formulation of plastisol, shape of the material, temperature and the like. In the a process for preparing a foamed polymer from the above-mentioned foamy polymer composition, the gelation time during the heating period in the presence of the functionalized particulate bicarbonate as the blowing agent is less than that in the presence of azodicarbonamide as the blowing agent (while all other components remain the same).

When foamy polymer composition comprises the functionalized particulate bicarbonate, the gelation time at temperature above glass transition temperature Tg and/or above melting temperature Tm of the polymer preferably is less than 180 seconds, such as less than 125 seconds.

Regarding a PVC plastisol, the resulting liquid PVC paste (plastisol) can be spread coated at a certain thickness on a surface (for example, a paper, in particular a silicone paper) and heated in an oven (e.g., Thermosol Werner Mathis) set at a curing (gelation) temperature (generally between 150°C and 210°C or 200°C, such as from 180°C to 210°C, or from 185°C to 210°C, or from 190°C to 210°C, or from 200°C to 210°C) for a time of 130 seconds or less, such as for a time of 120 seconds to 30 seconds, preferably for a time of 120 seconds to 60 seconds, more preferably for a time of 120 seconds to 90 seconds.

For some embodiments of a process for manufacturing a PVC polymer, when the foamy polymer composition is heated, the temperature suitable for liberating CO2 gas from the functionalized particulate sodium bicarbonate and melting a PVC polymer may be from 190 to 210°C, preferably from 200 to 210°C, during a gelation time from 90 seconds to 120 seconds at a temperature to provide a foamed PVC polymer.

When the foamy polymer composition is spread coated on a surface before heating and polymer melting, the foamed polymer may have an expansion ratio of at least 270, preferably at least 280, more preferably at least 300 and/or has a density of less than 0.6 g/cm³, preferably less than 0.55 g/cm³, more preferably at most 0.5 g/cm³. The expansion
ratio is calculated based on the ratio of a final thickness over an initial thickness of a spread coated layer of the foamable polymer composition as that layer gets heated in an oven.

Foamed polymeric products produced using the blowing agent and the above discussed foamable polymer composition can be made by processes that involve extrusion, calendering, injection molding, coating, expansion casting or rotational molding.

**FOAMED POLYMER**

A further aspect of the present invention relates to a foamable polymer, wherein the foamable polymer is obtainable from the foamable composition as described above.

In one embodiment of the invention, the foamable polymer is a flexible foamed polymer. In this embodiment, the foamed or extruded polymer is preferably flexible polyvinyl chloride and can be advantageously used

- for flooring applications;
- for production of wall paper foam based;
- as artificial leather; or
- as technical foam.

The functionalized particulate bicarbonate is preferably used as a blowing agent for foamed or extruded polymers (such as foamed PVC or polyurethanes; extruded PVC, polyolefins, polyamides), preferably used as an endothermic blowing agent. Non-limiting examples of polymers are polyvinyl chloride (PVC), polyurethanes, polyolefins (PO, PE, PP), styrenics (PS, ABS, ASA, SAN), engineering resins, polyphenylene oxide, polyamides, polyimides, and natural and synthetic rubber such as nitrile butadiene rubber (NBR) or chloroprene rubber (CR).

When the foaming agent contains the particulate bicarbonate without the polyoxyalkylene additive, the resulting foamable polymer composition has a higher viscosity than a foamable polymer composition containing a different chemical foaming agent: azodicarbonamide. Such high viscosity of the foamable polymer composition (with only particulate bicarbonate) results in a foamable polymer composition which is denser than a foamed polymer made with azodicarbonamide as foaming agent. The higher viscosity obtained with particulate non-functionalized bicarbonate used as the chemical foaming agent generally results in poor density (>0.6 g/ml) and low thickness (< 2 mm) of a PVC sheet made from the foamable composition.

On the other end, the addition of the polyoxyalkylene addition to functionalize particulate bicarbonate used in the chemical blowing agent can provide a foamable polymer
composition with a much reduced viscosity compared to a foamable polymer composition with non-functionalized particulate bicarbonate. An acceptable viscosity for the foamable composition may range from 9000 to 23000 mPa sec at 20°C at a shear rate of 3.14 s⁻¹ or range from 11000 to 21000 mPa sec at 20°C at a shear rate of 10.2 s⁻¹. The lower viscosity obtained with functionalized particulate bicarbonate as chemical blowing agent provides a PVC sheet made from the foamable composition with a good density (≤ 0.6 g/ml, preferably ≤ 0.5 g/ml) and a spread coat thickness ≥ 2 mm. With a reference thickness of the non-foamed material of 0.75 mm this corresponds to an expansion ratio of about >27().

When the foamable PVC polymer composition comprising the functionalized particulate bicarbonate is spread coated on a surface before heating and polymer melting, the foamed polymer may have an expansion ratio of at least 270, preferably at least 280, more preferably at least 300. The expansion ratio is calculated based on the ratio of a final thickness over an initial thickness of a spread coated layer of the foamable polymer composition as that layer gets heated in an oven. The expansion ratio may be from 270 up to 450, or from 280 up to 445, or from 290 up to 440, or from 300 up to 440.

In some embodiments, the foamed polymer is a PVC foam with a density of less than 0.65 g/cm³ or of less than 0.6 g/cm³, preferably less than 0.58 g/cm³, more preferably at most 0.55 g/cm³, most preferably at most 0.5 g/cm³ or at most 0.45 g/cm³. In some embodiments in which a PVC foam is made from a foamable PVC composition containing a functionalized particulate bicarbonate with at least one additive, the PVC foam has a density from 0.3 to 0.65 g/cm³, preferably a density from 0.33 to 0.58 g/cm³, more preferably a density from 0.33 to 0.5 g/cm³ or even a density from 0.33 to 0.45 g/cm³.

The additive in the functionalized particulate bicarbonate preferably comprises a polyoxyethylene compound (like BI3 155, polyethylene glycol e.g., PEG400, PEG4000, polypropylene glycol, e.g., PPG400, PPG425). Additional functionalized additive may be selected from the group consisting of beeswax, carnauba wax, glycerol mono stearate, poly(meth)acrylate, epoxidized soybean oil, arabic gum, linoleic acid, maltodextrin, polyvinyl alcohol, lauric acid, rosin acid or any derivatives thereof, starch, stearic acid, citric acid, leucine, and any combination of two or more thereof.

The present invention is based on the discovery of the effects of the additive mentioned in the present specification, for which the different variants of the method and/or of the product obtained by said variants of the method are described in more detail below.
ITEM 1. A chemical blowing agent for foaming a thermoplastic polymer precursor, for example PVC plastisol or a polymer resin in an extrusion process, said chemical blowing agent comprising a functionalized particulate bicarbonate, wherein said functionalized particulate bicarbonate contains at least one functionalization additive, and wherein said functionalization additive in the functionalized particulate bicarbonate comprises a polyoxyalkylene additive selected from the group consisting of polyoxyalkylenes, any derivatives thereof (such as esters), salts thereof, and any combination of two or more thereof.

ITEM 2. The chemical blowing agent according to ITEM 1, wherein the blowing agent does not contain any further blowing agent which is an exothermic blowing agent, or wherein the chemical blowing agent does not contain a compound which liberates nitrogen gas during heating, or wherein the chemical blowing agent does not contain a compound which liberates ammonia gas during heating.

ITEM 3. The chemical blowing agent according to ITEM 1 or 2, wherein the polyoxyalkylene additive comprises:

- a polyethylene oxide (PEO) polymer, preferably of a MW of 20,000 or more,
- a polyethylene glycol (PEG) polymer, preferably of a MW of less than 20,000, or a MW of 4000 or less, or a MW of 3600 or less, or a MW of 2000 or less, or a MW of 1000 or less, or a MW of 800 or less,
- a polypropylene oxide (PPO) polymer,
- a polypropylene glycol (PPG) polymer,
- a co-polymer or block polymer which is made with at least one monomer of propylene oxide or propylene glycol,
- a co-polymer or block polymer which is made with at least one monomer of ethylene oxide or ethylene glycol,
- a polyethylene glycol / polypropylene glycol (PEG/PPG) block co-polymer, such as the PEG/PPG/PEG triblock polymer, or any derivatives (such as esters) thereof,

wherein any of these polymers may be chemically functionalized or not.
ITEM 4. The chemical blowing agent according to any of ITEMS 1 to 3, wherein the polyoxyalkylene additive comprises a polyethylene glycol, preferably a polyethylene glycol with a molecular weight less than 1000 g/mol.

ITEM 5. The chemical blowing agent according to any of ITEMS 1 to 3, wherein the polyoxyalkylene additive comprises a polyoxyalkylene derivative which is an ester, such as a PEG or PPG ester.

ITEM 6. The chemical blowing agent according to any of ITEMS 1 to 3, wherein the polyoxyalkylene additive is a polyoxyalkylene derivative which is a chemically functionalized polyoxyalkylene, for example, a polyoxyalkylene bearing one or more reactive substituents, such as -OH, -NH₂, -NHR, -SH and -COOH, wherein R is a hydrocarbon residue, and which has been reacted with a functional chemical compound being reactive with the reactive substituent of the polyoxyalkylene.

ITEM 7. The chemical blowing agent according to any of ITEMS 1 to 6, wherein the functionalized particulate bicarbonate comprises at least one additional functionalization additive selected from
- one or more polymers other than the polyoxyalkylene additive;
- one or more amino acids, any derivative thereof, and salts thereof;
- one or more inorganic salts;
- one or more oils;
- one or more fats;
- one or more resin acids, any derivative thereof, and salts thereof;
- one or more fatty acids, any derivative thereof, and salts thereof;
- a carboxylic or polycarboxylic acid, derivative thereof (such as esters), or salts thereof;
- one or more soaps;
- one or more waxes; or
- any combinations thereof;
preferably selected from the group consisting of:
- one or more fatty acids, any derivative thereof, and salts thereof;
- a carboxylic or polycarboxylic acid, derivative thereof (such as esters), or salts thereof;
and
- one or more resin acids, any derivative thereof, and salts thereof.
ITEM 8. The chemical blowing agent according to any of ITEMS 1 to 7, wherein the functionalized particulate bicarbonate comprises at least one additional functionalization additive, said additional functionalization additive comprising:

- rosin; and/or
- a derivative of rosin acid, preferably a derivative of rosin acid selected from the group consisting of rosin acid esters, such as Ci-25 alkyl esters of rosin, glycerol rosin esters, pentaerythritol rosin esters, or combinations thereof, hydrogenated rosin acid, dimers of rosin acid, polymerized rosin, more preferably dihydroabietic acid, such as at least 50 wt% of dihydroabietic acid.

ITEM 9. The chemical blowing agent according to any of ITEMS 1 to 7, wherein the functionalized particulate bicarbonate comprises at least one additional functionalization additive, said additional functionalization additive comprising: abietic acid, dihydroabietic acid, a rosin acid ester, or mixtures thereof, preferably comprising abietic acid, dihydroabietic acid, or mixtures thereof.

ITEM 10. The chemical blowing agent according to any of ITEMS 1 to 9, wherein the functionalized particulate bicarbonate comprises at least 50% by weight and less than 100% by weight of the bicarbonate component, and from 50% or less to 0.02% by weight of at least one of said functionalization additive, preferably comprises at least 65% by weight and less than 100% by weight of the bicarbonate component, and from 35% or less to 0.02% by weight of at least one of said functionalization additive.

ITEM 11. The chemical blowing agent according to any of ITEMS 1 to 10 wherein the particles of the functionalized particulate bicarbonate have a particle size distribution of $D_{50}$ of more than 1 µm and at most 250 µm, preferably at most 100 µm, more preferably at most 60 µm, yet more preferably at most 40 µm, yet most preferably at most 25 µm.

ITEM 12. The chemical blowing agent according to any of ITEMS 1 to 10, wherein the particles of the functionalized particulate bicarbonate have a particle size distribution of $D_{50}$ of at most 1 µm.
ITEM 13. The chemical blowing agent according to any of ITEMS 1 to 12, wherein the functionalized particulate bicarbonate is obtained by at least one of the following processes:
- by spray-drying (also known as atomization), wherein the additive is dissolved in the bicarbonate-containing solution.
- by grinding or co-grinding (also known as milling or co-milling) with the additive(s) in emulsion or powder form;
- by spray coating and granulation within a fluidized bed,
- by spray agglomeration within a fluidized bed,
- by spray chilling (e.g., spray cooling, spray freezing),
- by roller compaction, and/or
- by extrusion, including simultaneous mixing/extrusion;
  optionally followed by subjecting the thus obtained functionalized particulate bicarbonate to milling to reduce its mean particle size; preferably obtained by co-grinding, extrusion, and/or spray coating with the additive(s); more preferably obtained by grinding and/or extrusion with the additive(s), or obtained by spray coating and/or extrusion with the additive(s).

ITEM 14. The chemical blowing agent according to any of ITEMS 1 to 13, further comprising an second compound which liberates CO2 upon heating, said second compound being selected from the group consisting of a carboxylic or polycarboxylic acid, derivative thereof (such as esters), or salts thereof, wherein said second compound optionally is functionalized with at least one additive which is different or the same as the one in the functionalized particulate bicarbonate, preferably the same additive, and wherein said second compound preferably is at least one of:
- fumaric acid,
- tartaric acid,
- citric acid, citrates (such as sodium hydrogen citrate, disodium citrate), or esters of citric acid; or combination thereof.

ITEM 15. The chemical blowing agent according to any of ITEMS 1 to 14, wherein the chemical blowing agent or the functionalized particulate bicarbonate further comprises silica, preferably amorphous silica, preferably amorphous precipitated silica.
ITEM 16. The chemical blowing agent according to any of ITEM 15, wherein the silica, is an amorphous silica having a high surface area of at least 180 m²/g, or of at least 200 m²/g, or of at least 400 m²/g, and/or a mean diameter of less than 20 microns, preferably of at most 15 microns.

ITEM 17. A foamable polymer composition, comprising a polymer, optionally a foaming stabilizer, and the chemical blowing agent according to any of ITEMS 1 to 16, wherein the polymer is a PVC, a polyurethane, a polyolefin, or a polyamide.

ITEM 18. The foamable polymer composition according to ITEM 17, further comprising an exothermic chemical blowing agent, preferably selected from the group consisting of ADCA (azodicarbonamide), OBSH (4,4'-oxybis(benzenesulfonyl hydrazide)), DNPT (dinitrosopentamethylene tetramine), PTSS (p-toluene semicarbazide), BSH (benzene-4-sulfonohydrazide), and TSH (toluene-4-sulfonohydrazide), 5-PT (5-phenyltetrazole), more preferably comprising ADCA.

ITEM 19. The foamable polymer composition according to ITEM 17 or 18, being a foamable PVC plastisol composition, wherein the polymer comprises a PVC resin.

ITEM 20. A foamable PVC plastisol composition, comprising a PVC resin, optionally a foaming stabilizer, and the chemical blowing agent according to any of ITEM 1 to 16, which preferably does not contain azodicarbonamide, benzenesulfonylhydrazide, and/or p-toluenesulfonhydrazide.

ITEM 21. A process for manufacturing a polymer, comprising heating the foamable polymer composition according to ITEM 17 which comprises the functionalized particulate bicarbonate at a temperature suitable for liberating CO₂ gas and melting the polymer during a gelation time at a temperature above glass transition temperature Tg and/or above melting temperature Tm of the polymer which is less than 130 seconds.

ITEM 22. A foamed polymer obtained by the process of ITEM 21, wherein the polymer is a PVC, a polyurethane, a polyolefin, or a polyamide.
ITEM 23. A process for manufacturing a foamed PVC polymer, comprising heating the foamable PVC polymer composition according to ITEM 20 or 21 which comprises the functionalized particulate bicarbonate and the PVC resin at a temperature suitable for liberating CO₂ gas and melting the PVC resin during a gelation time from 90 seconds to 120 seconds at a temperature from 190 to 210°C, preferably from 200 to 210°C, to provide a foamed PVC polymer, wherein the foamable polymer composition preferably is spread coated on a surface before heating and melting, and wherein the foamed polymer has an expansion ratio of at least 270, preferably at least 280, more preferably at least 300 and/or has a density of less than 0.6 g/cm³, preferably less than 0.55 g/cm³, more preferably at most 0.5 g/cm³.

ITEM 24. A process for manufacturing a polymer, comprising heating the foamable PVC plastisol composition according to ITEM 20 or 21 which comprises the functionalized particulate bicarbonate at a temperature suitable for liberating CO₂ gas and melting the PVC resin during a gelation time, wherein said gelation time with the chemical blowing agent comprising the functionalized particulate is less than the gelation time that would be obtained with a chemical blowing agent liberating N₂, all other components in the foamable PVC plastisol composition being the same.

ITEM 25. A foamed PVC obtained by the process of ITEMS 23 or 24.

The following examples are given by way of non-limiting illustration of the present invention, and variations thereof that are readily accessible to a person skilled in the art.

EXAMPLES

Co-Milling

Milling was performed continuously in an UltraRotor III instrument from Jackering, equipped with a classifier. Sodium Bicarbonate particles were loaded at the bottom of the mill from 10 to 300 kg/h under air injection at room temperature. The functionalization additive was added at the mid level of the mill in order to reach the desired weight content of functionalization additive in the mix. Mill rotary speed and classifier rotary speed were selected to reach the desired particle size distribution.
**Extrusion-Mixing**

The extrusion-mixing process was performed continuously in a UCP25 instrument from HASLER Group which had an inlet point and paddles. Sodium Bicarbonate particles were loaded at the UCP25 inlet point from 1 to 2 kg/h. The functionalization additive was loaded a few centimeters after the main inlet point, from 0.1 to 1 kg/h. Rotary speed of the paddles was 50 rpm. Instrument temperature was kept at room temperature.

**Milling after Extrusion-Mixing**

The functionalized sodium bicarbonate product exiting the extrusion-mixing process was continuously injected at the top of a UPZ100 mill from Hosokawa Alpine at a loading rate of from 0.5 to 10 kg/h. The mill has a rotor speed selected between 10,000 and 17,000 rpm and the mill was operated at room temperature.

**Fluid Bed Spray Coating (= FB Coating + milling)**

Spray coating was performed in a batch fluidized bed WFP-mini instrument from DMR Prozesstechnologie which comprised a fluidization chamber, means to load particles, means to inject the functionalization additive generally in liquid form, and means to inject a fluidization gas at the bottom of that chamber. Sodium Bicarbonate particles were initially loaded from 100 g to 1 kg in the fluidization chamber. Fluidization air was heated from 10 to 100 °C, and its flowrate was between 10 and 40 m³/h. A liquid containing the functionalization additive was sprayed from the chamber bottom at a temperature between 20 and 90 °C. The liquid was generally a solution of the functionalization additive in a solvent, preferably an aqueous solution with a concentration between 1 and 80 wt%. The liquid was sprayed under a flowrate of up to 20 g/min in order to evaporate the solvent (e.g., water) and coat the functionalization additive (generally by way of crystallization) onto the sodium bicarbonate particles. The content of the coated functionalization additive was between 1 and 70 wt%.

The product obtained after spray coating was generally milled. The spray coated particles that exited the fluid bed coating process were injected at the top of a UPZ100 mill.
from Hosokawa Alpine at a loading rate of from 0.5 to 10 kg/h and the mill had a rotor speed selected between 10,000 and 17,000 rpm and was operated at room temperature.

Preparation of PVC plastisol

A method for making a PVC plastisol may generally follow the following steps:
- the weight of the PVC resin is measured in a 2-liter stainless steel pot and the rest of the formula is weighed apart:
- The stainless steel pot is placed under a hydraulic mixer (Pendraulik) with a deflocculating shear blade (rotor blade diameter 70 mm)
- Stirring at 250 rev / min during addition of the mineral filler, plasticizer and blowing agent
- Stirring at 4200 rev / min for 45 seconds, once the powders are suspended (temperature about 40-50 °C, if time higher agitation, T may rise to 60-70 °C)
- Vacuum preparation to strip air bubbles. When the level of preparation goes up into the beaker, the beaker may be tapped on the bench to remove bubbles faster.
- When the level of preparation is stable in the presence of the vacuum, count 5 minutes before cutting vacuum and then end air stripping.
- To avoid foaming, optionally adding a foam stabilizer (such as from BYK provider)
- temperature setting in a Werner Mathis oven at a preset temperature, usually T = 190, 200 or 210 °C
- A paper (silicone paper in examples 1-8 and regular paper (from Claire Fontaine) in examples 9-13) is installed and heated for 10 seconds to remove residual water from the paper
- Plastisol mixture is spread on the paper in a thickness of 750 microns
- The PVC plastisol is heated up to 2 minutes for gelation (usually 60s, 90s, 120s),
- Wait until the plastisol is cooled for remove it from the paper.

The following chemicals were used in the examples:

SolVin® 367 NF  PVC resin  oflnovyn
SolVin® 266 SC  PVC resin  of Inovyn
Durcal® 15     CaC0₃  of Omya
Palatinol®     (DINP) di-isononyl of BASF
Example 1 (not in accordance with invention)

The PVC Plastisol composition of Example 1 was prepared by using azodicarbonamide as the blowing agent. The other components of the plastisol of Example 1 were at least one polyvinyl chloride polymer, and optionally one or more polymer additives such as an inorganic filler.

The specific components and amounts as mentioned in Table 1 were chosen. In each example, the components were dispersed on a high speed mixer and then mixed and de-aerated under vacuum.

The resulting liquid PVC paste (plastisol) of Example 1 was spread coated at a thickness of about 0.75 mm on silicone paper and heated in a Thermosol Werner Mathis oven set at 185 °C for 120 seconds.

The surface and structure of the foam was assessed visually. The expansion rate was the ratio of the difference between the final thickness of the foamed product after Werner
Mathis oven heating and the initial thickness of the spread coated layer (here about 0.75 mm) before it passed the Werner Mathis oven heating, over the initial thickness. The expansion ratio was calculated as the ratio of the final thickness of the foamed product over the initial thickness of the spread coated layer (here about 0.75 mm). The results are provided in TABLE 1, where "PCR" means parts by weight of PVC resin.

### TABLE 1

<table>
<thead>
<tr>
<th>Example</th>
<th>1 PLASTISOL BASE AZO PCR</th>
</tr>
</thead>
<tbody>
<tr>
<td>PVC resin</td>
<td>SolVin 367 NF 89.9</td>
</tr>
<tr>
<td></td>
<td>SolVin 266 SC 10.1</td>
</tr>
<tr>
<td>Inorganic filler</td>
<td>Durcal 15 (CaCO₃) 80.1</td>
</tr>
<tr>
<td>Plastisizer</td>
<td>Foam stabilizer (DINP) 58.1</td>
</tr>
<tr>
<td>Wetting and dispersing agent of TiO₂ and ZnO</td>
<td>Disperplast 1150 0.3</td>
</tr>
<tr>
<td>Viscosity Modifiers</td>
<td>Viscobyk 4040 3.2</td>
</tr>
<tr>
<td>Prevent the foam release agent</td>
<td>BYK 3155 0.3</td>
</tr>
<tr>
<td>Foam stabilizer</td>
<td>BYK 8070 -</td>
</tr>
<tr>
<td></td>
<td>TiO₂ 2.3</td>
</tr>
<tr>
<td>Blowing agent</td>
<td>Azodicarbonamide 2.2</td>
</tr>
<tr>
<td></td>
<td>ZnO 1</td>
</tr>
</tbody>
</table>

| Conditions/Results | Pre-gelification phase | 30 sec at 90°C |
|                    | Gelification phase     | 2 min at 185°C |
|                    | Coating appearance     | Very smooth |
|                    | Thickness (in mm)      | 2.4 |
|                    | Expansion rate (%)     | 220 |
|                    | Expansion ratio (%)    | 320 |

Examples 2-5 (not in accordance with invention)

For Examples 3-5, PVC Plastisol compositions were prepared by using a non-functionalized sodium bicarbonate in powder form as the blowing agent. The non-functionalized sodium bicarbonate in powder form was mixed in the presence at least one
plasticizer, at least one polyvinyl chloride polymer, and optionally one or more polymer additives such as an inorganic filler. A PVC Plastisol composition of Example 2 was prepared by using azodicarbonamide as the blowing agent. The other components of the plastisol of Example 2 were the same as those used in Examples 3-5. The specific components and amounts as mentioned in TABLE 2 were chosen. "PCR" in TABLE 2 means parts by weight of resin. In each example 2-5, the components were dispersed on a high speed mixer and then mixed and de-aerated under vacuum.

The resulting liquid PVC pastes (plastisols) of Examples 2-5 were in each case spread coated at a thickness of about 0.75 mm on silicone paper and heated in a Thermosol Werner Mathis oven set with a gelation period at 185 °C for 120 seconds.

Examples 3 to 5 with non-functionalized bicarbonates as foaming agent gave poor expansion results compared to the Example 2 using azodicarbonamide as the blowing agent. After gelling, PVC plastisols surfaces of Examples 3 to 5 had a very granular appearance which was a sign of surface coalescence of gas bubbles in the polymer melt.

| TABLE 2 |
|---|---|---|---|---|
| Examples | 2 | 3 | 4 | 5 |
| | A | B | C | D |
| PLASTISOL BASE AZO | Bicar 0/6P | Bica 0/50 | Bicar 0/50 |
| PCR | PCR | PCR | PCR |
| PVC resin | PVC resin | 100 | 100 | 100 |
| Inorganic filler | CaCO₃ | 40 | 40 | 40 | 40 |
| Plasticizer | Palatinol N (DINP) | 65 | 65 | 65 | 65 |
| | MB10 | | | | 20 |
| Blowing Agent | Azodicarbonamide + Kicker (6) | 4.5 | 1.1 | 1.1 | 1.1 |
| | Bicar 0/50 crushed (7) | X | 8 | 8 |
| | Bicar 0/6P (2) | X | 6 |
| Result | Coating appearance | Very smooth | Very granular | Very granular | Very granular |

(1) Bicar 06/P trade product of Solvay
(2) Bicar 0/50 crushed in Alpine 01 - dio = 3.4 µm ds = 17.8 µm dₛ₅ = 39.6 µm
Example 6 (not in accordance with invention)

The PVC Plastisol composition of Example 6 was prepared by using azodicarbonamide as the blowing agent. The other components of the plastisol of Example 6 were at least one plasticizer, at least one polyvinyl chloride polymer, and optionally one or more polymer additives such as an inorganic filler. The specific components and amounts as mentioned in TABLE 3 are chosen.

In each example, the components were dispersed on a high speed mixer and then mixed and de-aerated under vacuum.

The resulting liquid PVC paste (plastisol) of Example 6 was spread coated at a thickness of about 0.75 mm on silicone paper and heated in a Thermosol Werner Mathis oven set at 185°C for 120 seconds for Example 6.

The surface and structure of the foam was assessed visually. The expansion rate was the ratio of the difference between the final thickness of the foamed product after Werner Mathis oven heating and the initial thickness of the spread coated layer (here about 0.75 mm) before it passed the Werner Mathis oven heating, over the initial thickness. The expansion ratio was calculated as the ratio of the final thickness of the foamed product over the initial thickness of the spread coated layer (here about 0.75 mm). The results are provided in TABLE 3, where "per" means parts by weight of PVC resin.
TABLE 3

<table>
<thead>
<tr>
<th>Example 7 (not in accordance with invention)</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>Blowing agent = azodicarbonamide</td>
<td></td>
</tr>
</tbody>
</table>

The PVC Plastisol compositions of Example 7 were prepared in a Dispermat apparatus by using azodicarbonamide as the blowing agent. The azodicarbonamide product used was Genitron SCE (Lanxess) which consisted of 75 wt% azodicarbonamide and 25 wt% kicker (zinc oxide/zinc stearate). The components of the plastisol were dispersed on a high speed mixer and then mixed and de-aerated under vacuum. The resulting liquid PVC pastes (plastisols) of Example 7 are in each case spread coated at a thickness of about 0.75 mm on a piece of paper (Claire Fontaine) and heated in a Thermosol Werner Mathis oven set at 200 or 210°C for 90 or 120 seconds.
The expansion rate was the ratio of the difference between the final thickness of the foamed product after Werner Mathis oven heating and the initial thickness of the spread coated layer (here about 0.75 mm) before it passed the Werner Mathis oven heating, over the initial thickness. The expansion ratio was calculated as the ratio of the final thickness of the foamed product over the initial thickness of the spread coated layer (here about 0.75 mm).

The specific components and amounts in each plastisol, the oven temperature, the time for gelation, the thickness of the foamed product after oven heating, the expansion rate, the expansion ratio, and the density of the PVC foams are provided in TABLE 4 where “pcr” means parts by weight of PVC resin.

<table>
<thead>
<tr>
<th>TABLE 4: Azodicarbonamide as blowing agent</th>
</tr>
</thead>
<tbody>
<tr>
<td>Example</td>
</tr>
<tr>
<td>Type of blowing agent</td>
</tr>
<tr>
<td>Plastisol Ex.</td>
</tr>
<tr>
<td>PVC resin 367 NF</td>
</tr>
<tr>
<td>Filler CaCO₃</td>
</tr>
<tr>
<td>Plastisizer Palatinol N</td>
</tr>
<tr>
<td>Defoamer BYK3155</td>
</tr>
<tr>
<td>Blowing agent Genitron SCE (incl. 0.25% zinc oxide/stearate)</td>
</tr>
<tr>
<td>Gelation temp (°C)</td>
</tr>
<tr>
<td>Gelation Time (sec.)</td>
</tr>
<tr>
<td>Thickness (mm)</td>
</tr>
<tr>
<td>Expansion rate (%)</td>
</tr>
<tr>
<td>Expansion ratio (%)</td>
</tr>
<tr>
<td>Density (g/cm³)</td>
</tr>
</tbody>
</table>

Example 8 (not in accordance with invention)

With Functionalization Additive = Beeswax

One sample of functionalized sodium bicarbonate was prepared by spray coating sodium bicarbonate particles in a fluidized bed with beeswax (not a polyoxyalkylene) and then milling. The functionalized sodium bicarbonate sample is identified as Example 8A (20 wt% beeswax).

For Example 8, PVC Plastisol compositions were prepared in a Dispermat apparatus. In each example, the components of the plastisols were dispersed on a high speed mixer and then mixed and de-aerated under vacuum. The resulting liquid PVC pastes (plastisols) are in each case spread coated at a thickness of about 0.75 mm on a piece of paper (Claire Fontaine)
and heated in a Thermosol Werner Mathis oven set at 200 or 210 oC for 90 or 120 seconds. The expansion ratio is calculated as the ratio of the thickness of the foamed product after Werner Mathis oven heating over the initial thickness of the spread coated layer (here about 0.75 mm) before it passed the Werner Mathis oven heating. The specific components and amounts in each plastisol, the oven temperature, the time for gelation, the thickness of the foamed product after oven heating, the expansion ratio, and the density of the PVC foams are provided in TABLE 5, where "per" means parts by weight of PVC resin.

TABLE 5 : Functionalization Additive = Beeswax

<table>
<thead>
<tr>
<th>Example</th>
<th>8</th>
</tr>
</thead>
<tbody>
<tr>
<td>Functionalyzed Bicarbonate</td>
<td>Ex. 8A</td>
</tr>
<tr>
<td>Funct. Method</td>
<td>Spray Coating with fluidized bed / milling</td>
</tr>
<tr>
<td>Funct. Additive</td>
<td>20 wt% Beeswax</td>
</tr>
<tr>
<td>D10 / D50 / D90 (μm)</td>
<td>4.4 / 21.65</td>
</tr>
<tr>
<td>PVC resin</td>
<td>367 NF (pcr)</td>
</tr>
<tr>
<td>Plastisol Ex.</td>
<td>8 p1</td>
</tr>
<tr>
<td>Filler</td>
<td>CaCO3 (pcr)</td>
</tr>
<tr>
<td>120</td>
<td>120</td>
</tr>
<tr>
<td>Plastisizer</td>
<td>Palatinol N (pcr)</td>
</tr>
<tr>
<td>80</td>
<td>80</td>
</tr>
<tr>
<td>Defoamer</td>
<td>BYK3155 (pcr)</td>
</tr>
<tr>
<td>3</td>
<td>3</td>
</tr>
<tr>
<td>Blowing agent</td>
<td>Func. Bicar. (pcr)</td>
</tr>
<tr>
<td>8</td>
<td>8</td>
</tr>
<tr>
<td>Gelation temp (°C)</td>
<td>200</td>
</tr>
<tr>
<td>Gelation Time (sec.)</td>
<td>90</td>
</tr>
<tr>
<td>Thickness (mm)</td>
<td>2.11</td>
</tr>
<tr>
<td>Expansion ratio (%)</td>
<td>280</td>
</tr>
<tr>
<td>Density (g/cm³)</td>
<td>0.53</td>
</tr>
</tbody>
</table>

Examples 9-10 (in accordance with invention)

Functionalization Polyoxyalkylene Additive = Polyethylene Glycol (PEG400)

Two samples of functionalized sodium bicarbonate were prepared by extrusion of sodium bicarbonate particles with a polyethylene glycol of low molecular weight (<1000 g/mol): PEG400 as the functionalization additive. The functionalized sodium bicarbonate samples are identified as Example 9A (20 wt% PEG400) and Example 10A (10 wt% PEG400).

For Examples 9-10, the same method of preparation of the PVC Plastisol compositions and preparation of PVC foams were used as described for Example 8. The expansion ratio was calculated as the ratio of the thickness of the foamed product after Werner Mathis oven heating over the initial thickness of the spread coated layer (here about 0.75 mm) before it
passed the Werner Mathis oven heating. The specific components and amounts in each plastisol, the oven temperature, the time for gelation, the thickness of the foamed product after oven heating, the expansion ratio and the density of the PVC foams are provided in TABLE 6, where "per" means parts by weight of PVC resin.

### TABLE 6: Functionalization Polyoxyalkylene Additive = Polyethylene Glycol (PEG400)

<table>
<thead>
<tr>
<th>Example</th>
<th>9</th>
<th>10</th>
</tr>
</thead>
<tbody>
<tr>
<td>Functionalized Bicarbonate</td>
<td>Ex. 9A</td>
<td>Ex. 10A</td>
</tr>
<tr>
<td>Funct. Method</td>
<td>extrusion</td>
<td>extrusion</td>
</tr>
<tr>
<td>Funct. additive</td>
<td>20 wt% PEG400</td>
<td>10 wt% PEG400</td>
</tr>
<tr>
<td>D10/D50/D90 (μm)</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>PVC resin</td>
<td>367 NF (pcr)</td>
<td>100</td>
</tr>
<tr>
<td>Filler</td>
<td>CaCO₃ (pcr)</td>
<td>120</td>
</tr>
<tr>
<td>Plastisizer</td>
<td>Palatinol N (pcr)</td>
<td>80</td>
</tr>
<tr>
<td>Defoamer</td>
<td>BYK3155 (pcr)</td>
<td>-</td>
</tr>
<tr>
<td>Blowing agent</td>
<td>Func. Bicar. (pcr)</td>
<td>10</td>
</tr>
<tr>
<td>Gelation temp (°C)</td>
<td>200</td>
<td>200</td>
</tr>
<tr>
<td>Gelation Time (sec.)</td>
<td>90</td>
<td>90</td>
</tr>
<tr>
<td>Thickness (mm)</td>
<td>2.52</td>
<td>2.14</td>
</tr>
<tr>
<td>Expansion ratio (%)</td>
<td>340</td>
<td>340</td>
</tr>
<tr>
<td>Density (g/cm³)</td>
<td>0.44</td>
<td>0.44</td>
</tr>
</tbody>
</table>

**Examples 11-15 (in accordance with invention)**

**Functionalization Polyoxyalkylene Additive = Polyethylene Glycol (PEG400)**

Three samples of functionalized sodium bicarbonate were prepared by extrusion of sodium bicarbonate particles with a polyethylene glycol of high molecular weight (>1000): PEG 4000, and then milling. Two other sample of functionalized sodium bicarbonate were prepared by spray coating of sodium bicarbonate particles in a fluidized bed with PEG4000, and then milling. The functionalized sodium bicarbonate samples are identified as Example 11A (extruded with 7 wt% PEG4000), Example 12A (extruded with 10 wt% PEG4000), Example 13A (extruded with 3 wt% PEG4000), Example 14A (spray coated with 10 wt% PEG4000), and Example 15A (spray coated with 20 wt% PEG4000).

For Examples 11-15, the same method of preparation of the PVC Plastisol compositions and preparation of PVC foams were used as described for Example 8. The expansion ratio was calculated as the ratio of the thickness of the foamed product after...
Werner Mathis oven heating over the initial thickness of the spread coated layer (here about 0.75 mm) before it passed the Werner Mathis oven heating. The specific components and amounts in each plastisol, the oven temperature, the time for gelation, the thickness of the foamed product after oven heating, the expansion ratio and the density of the PVC foams are provided in TABLE 7, where "per" means parts by weight of resin.

**TABLE 7 : Functionalization Polyoxyalkylene Additive = Polyethylene Glycol (PEG4000)**

<table>
<thead>
<tr>
<th>Example</th>
<th>11</th>
<th>12</th>
</tr>
</thead>
<tbody>
<tr>
<td>Functionized Bicarbonate</td>
<td>Ex. 11A</td>
<td>Ex. 12A</td>
</tr>
<tr>
<td>Funct. Method</td>
<td>Extrusion / milling</td>
<td>Extrusion/ milling</td>
</tr>
<tr>
<td>Funct. additive</td>
<td>7 wt% PEG4000</td>
<td>10 wt% PEG4000</td>
</tr>
<tr>
<td>D10 / D50 / D90 (μm)</td>
<td>4.9 / 22 / 57</td>
<td>6.9 / 32 / 81</td>
</tr>
<tr>
<td>Plastisol Ex.</td>
<td>11 p1 11 p2 11 p3 11 p4</td>
<td>12 p1 12 p2 12 p3</td>
</tr>
<tr>
<td>PVC resin</td>
<td>367 NF (pct)</td>
<td>100 100 100 100</td>
</tr>
<tr>
<td>Filler</td>
<td>CaCO₃ (pct)</td>
<td>120 120 120 120</td>
</tr>
<tr>
<td>Plastisizer</td>
<td>Palatinol N (pct)</td>
<td>80 80 80 80</td>
</tr>
<tr>
<td>Defoamer</td>
<td>BYK3155 (pct)</td>
<td>3 3 3 3</td>
</tr>
<tr>
<td>Blowing agent</td>
<td>Func. Bicar. (pct)</td>
<td>8 8 8 8</td>
</tr>
<tr>
<td>Gelation temp (°C)</td>
<td>200 210 210 210</td>
<td>200 210 210</td>
</tr>
<tr>
<td>Gelation Time (sec.)</td>
<td>2.16 2.29 2.97 2.39</td>
<td>2.46 2.89 2.19</td>
</tr>
<tr>
<td>Thickness (mm)</td>
<td>0.54 0.50 0.37 0.43</td>
<td>0.43 0.37 0.48</td>
</tr>
<tr>
<td>Expansion ratio (%)</td>
<td>290 400 320 330</td>
<td>390 290</td>
</tr>
<tr>
<td>Density (g/cm³)</td>
<td>0.54 0.50 0.37 0.43</td>
<td>0.43 0.37 0.48</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Example</th>
<th>13</th>
<th>14</th>
</tr>
</thead>
<tbody>
<tr>
<td>Functionized Bicarbonate</td>
<td>Ex. 13A</td>
<td>Ex. 14A</td>
</tr>
<tr>
<td>Funct. Method</td>
<td>Extrusion/ milling</td>
<td>Spray Coating/ milling</td>
</tr>
<tr>
<td>Funct. additive</td>
<td>3 wt% PEG4000</td>
<td>10 wt% PEG4000</td>
</tr>
<tr>
<td>D10 / D50 / D90 (μm)</td>
<td>5.2 / 25 / 61</td>
<td>5.6 / 26 / 65</td>
</tr>
<tr>
<td>Plastisol Ex.</td>
<td>13 p1 13 p2 13 p3 14 p1</td>
<td>14 p2 14 p3 14 p4</td>
</tr>
<tr>
<td>PVC resin</td>
<td>367 NF (pct)</td>
<td>100 100 100 100</td>
</tr>
<tr>
<td>Filler</td>
<td>CaCO₃ (pct)</td>
<td>120 120 120 120</td>
</tr>
<tr>
<td>Plastisizer</td>
<td>Palatinol N (pct)</td>
<td>80 80 80 80</td>
</tr>
<tr>
<td>Defoamer</td>
<td>BYK3155 (pct)</td>
<td>3 3 3 3</td>
</tr>
<tr>
<td>Blowing agent</td>
<td>Func. Bicar. (pct)</td>
<td>8 8 8 8</td>
</tr>
<tr>
<td>Gelation temp (°C)</td>
<td>200 210 210 210</td>
<td>210 210 210</td>
</tr>
</tbody>
</table>
Examples 16-17 (in accordance with invention)

**Functionalization** Polyoxyalkylene Additive = BYK 3155

Two samples of functionalized sodium bicarbonate were prepared by extrusion of sodium bicarbonate particles with a polyoxyalkylene compound as the functionalization additive. The polyoxyalkylene compound is a product sold by BYK under the brandname BYK 3155 which may be used as an air release agent. The functionalized sodium bicarbonate samples are identified as Example 16A (10 wt% BYK 3155) and Example 17A (20 wt% BYK 3155).

For Examples 16-17, the same method of preparation of the PVC Plastisol compositions and preparation of PVC foams were used as described for Example 8. The expansion ratio was calculated as the ratio of the thickness of the foamed product after Werner Mathis oven heating over the initial thickness of the spread coated layer (here about 0.75 mm) before it passed the Werner Mathis oven heating. The specific components and
amounts in each plastisol, the oven temperature, the time for gelation, the thickness of the
foamed product after oven heating, the expansion ratio and the density of the PVC foams are
provided in TABLE 8, where "per" means parts by weight of PVC resin.

### TABLE 8 : Functionalization Polyoxyalkylene Additive = BYK 3155

<table>
<thead>
<tr>
<th>Example</th>
<th>16</th>
<th>17</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Ex. 16A</td>
<td>Ex. 17A</td>
</tr>
<tr>
<td>Functionalized Bicarbonate</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Funct. Method</td>
<td>extrusion</td>
<td>extrusion</td>
</tr>
<tr>
<td>Funct. additive</td>
<td>10 wt% BYK3155</td>
<td>20 wt% BYK3155</td>
</tr>
<tr>
<td>PVC resin</td>
<td>367 NF (pcr)</td>
<td>100</td>
</tr>
<tr>
<td>Filler</td>
<td>CaCO₃ (pcr)</td>
<td>120</td>
</tr>
<tr>
<td>Plastisizer</td>
<td>Palatinol N (pcr)</td>
<td>80</td>
</tr>
<tr>
<td>Defoamer</td>
<td>BYK3155 (pcr)</td>
<td>-</td>
</tr>
<tr>
<td>Blowing agent</td>
<td>Func. Bicar. (pcr)</td>
<td>9</td>
</tr>
<tr>
<td>Gelation temp (°C)</td>
<td>200</td>
<td>200</td>
</tr>
<tr>
<td>Gelation Time (sec.)</td>
<td>90</td>
<td>90</td>
</tr>
<tr>
<td>Thickness (mm)</td>
<td>2.03</td>
<td>2.06</td>
</tr>
<tr>
<td>Expansion ratio (%)</td>
<td>270</td>
<td>270</td>
</tr>
<tr>
<td>Density (g/cm³)</td>
<td>0.52</td>
<td>0.47</td>
</tr>
</tbody>
</table>

**Examples 18-19 (in accordance with invention)**

**With Polyoxyalkylene Additive = Polyethylene Glycol (PEG400)**

**With Additional Functionalization Additive = Rosin acid**

Two samples of functionalized sodium bicarbonate were prepared by applying two
different functionalization additives (PEG400 as the polyoxyalkylene additive, rosin acid as
additional functionalization additive) added sequentially using two different methods of
functionalization (co-grinding, extrusion). First the sodium bicarbonate particles were
functionalized by co-grinding them with rosin acid as a first functionalization additive, and
then the first-functionalized particles were functionalized by extruding them with PEG400 as
a second functionalization additive to form second-functionalized particles. The samples are
identified as Example 18A (9 wt% rosin acid, 10 wt% PEG400) and Example 19A (8 wt% rosin acid, 20 wt% PEG400).

For Examples 18-19, the same method of preparation of the PVC Plastisol
compositions and preparation of PVC foams were used as described for Example 8. The
expansion ratio was calculated as the ratio of the thickness of the foamed product after Werner Mathis oven heating over the initial thickness of the spread coated layer (here about 0.75 mm) before it passed the Werner Mathis oven heating. The specific components and amounts in each plastisol, the oven temperature, the time for gelation, the thickness of the foamed product after oven heating, the expansion ratio and the density of the PVC foams are provided in TABLE 9 where "per" means parts by weight of PVC resin.

### TABLE 9: Two Functionalization Additives = Rosin acid (additional additive), PEG400 (polyoxyalkylene additive) added sequentially

<table>
<thead>
<tr>
<th>Example</th>
<th>18</th>
<th>19</th>
</tr>
</thead>
<tbody>
<tr>
<td>Functionalized Bicarbonate</td>
<td>Ex. 18A</td>
<td>Ex. 19A</td>
</tr>
<tr>
<td>1st Funct. Method</td>
<td>co-milling</td>
<td>co-milling</td>
</tr>
<tr>
<td>1st Funct. additive</td>
<td>10 wt% rosin acid</td>
<td>10 wt% rosin acid</td>
</tr>
<tr>
<td>2nd Funct. Method</td>
<td>extrusion</td>
<td>extrusion</td>
</tr>
<tr>
<td>2nd Funct. additive</td>
<td>10 wt% PEG400</td>
<td>20 wt% PEG400</td>
</tr>
<tr>
<td>Resulting Func. Bicar Composition</td>
<td>81 wt% Na bicarbonate / 9 wt% rosin acid / 10 wt% PEG400</td>
<td>72 wt% Na bicarbonate / 8 wt% rosin acid / 20 wt% PEG400</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Plastisol Ex.</th>
<th>18 p1</th>
<th>18 p2</th>
<th>18 p3</th>
<th>18 p4</th>
<th>19 p1</th>
<th>19 p2</th>
<th>19 p3</th>
</tr>
</thead>
<tbody>
<tr>
<td>PVC resin</td>
<td>367 NF (pcr)</td>
<td>100</td>
<td>100</td>
<td>100</td>
<td>100</td>
<td>100</td>
<td>100</td>
</tr>
<tr>
<td>Filler</td>
<td>CaCO₃ (pcr)</td>
<td>120</td>
<td>120</td>
<td>120</td>
<td>120</td>
<td>120</td>
<td>120</td>
</tr>
<tr>
<td>Plastisizer</td>
<td>Palatinol N (pcr)</td>
<td>80</td>
<td>80</td>
<td>80</td>
<td>80</td>
<td>80</td>
<td>80</td>
</tr>
<tr>
<td>Defoamer</td>
<td>BYK3155 (pcr)</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Blowing agent</td>
<td>Func. Bicar. (pcr)</td>
<td>9</td>
<td>9</td>
<td>9</td>
<td>9</td>
<td>10</td>
<td>10</td>
</tr>
<tr>
<td>Gelation temp (°C)</td>
<td>200</td>
<td>200</td>
<td>210</td>
<td>210</td>
<td>200</td>
<td>210</td>
<td>210</td>
</tr>
<tr>
<td>Gelation Time (sec.)</td>
<td>90</td>
<td>120</td>
<td>90</td>
<td>120</td>
<td>120</td>
<td>90</td>
<td>120</td>
</tr>
<tr>
<td>Thickness (mm)</td>
<td>2.03</td>
<td>2.06</td>
<td>2.30</td>
<td>2.08</td>
<td>2.43</td>
<td>2.60</td>
<td>2.45</td>
</tr>
<tr>
<td>Expansion ratio (%)</td>
<td>270</td>
<td>270</td>
<td>310</td>
<td>280</td>
<td>320</td>
<td>350</td>
<td>330</td>
</tr>
<tr>
<td>Density (g/cm³)</td>
<td>0.52</td>
<td>0.54</td>
<td>0.47</td>
<td>0.51</td>
<td>0.45</td>
<td>0.41</td>
<td>0.44</td>
</tr>
</tbody>
</table>

Examples 20-21 (in accordance with invention)

**With Polyoxyalkylene Additive = Polyethylene Glycol (PEG400)**

**With Additional Functionalization Additive = Stearic acid**

Two samples of functionalized sodium bicarbonate were prepared by applying two different functionalization additives (stearic acid, PEG400) added sequentially using two different methods of functionalization (co-grinding, extrusion). First the sodium bicarbonate
particles were functionalized by co-grinding them with stearic acid as a first functionalization additive, and then the first-functionalized particles were functionalized by extruding them with PEG400 as a second functionalization additive to form second-functionalized particles. The samples are identified as Example 20A (3.2 wt% stearic acid, 10 wt% PEG400) and Example 21A (2.8 wt% stearic acid, 20 wt% PEG400).

For Examples 20-21, the same method of preparation of the PVC Plastisol compositions and preparation of PVC foams were used as described for Example 8. The expansion ratio was calculated as the ratio of the thickness of the foamed product after Werner Mathis oven heating over the initial thickness of the spread coated layer (here about 0.75 mm) before it passed the Werner Mathis oven heating. The specific components and amounts in each plastisol, the oven temperature, the time for gelation, the thickness of the foamed product after oven heating, the expansion ratio, and the density of the PVC foams are provided in TABLE 10, where “per” means parts by weight of PVC resin.

### TABLE 10: Two Functionalization Additives = stearic acid (additional additive), PEG400 (polyoxyalkylene additive) added sequentially

<table>
<thead>
<tr>
<th>Example</th>
<th>20</th>
</tr>
</thead>
<tbody>
<tr>
<td>Functionalized Bicarbonate</td>
<td>Ex. 20A</td>
</tr>
<tr>
<td>1st Funct. Method</td>
<td>co-milling</td>
</tr>
<tr>
<td>1st Funct. additive</td>
<td>3.5 wt% stearic acid</td>
</tr>
<tr>
<td>2nd Funct. Method</td>
<td>extrusion</td>
</tr>
<tr>
<td>2nd Funct. additive</td>
<td>10 wt% PEG400</td>
</tr>
<tr>
<td>Resulting Func. Bicar Composition</td>
<td>86.9 wt% Na bicarbonate / 3.2 wt% stearic acid / 10 wt% PEG400</td>
</tr>
<tr>
<td>Plastisol Ex.</td>
<td>20 p1 20 p2 20 p3 20 p4 20 p5 20 p6 20 p7</td>
</tr>
<tr>
<td>PVC resin</td>
<td>367 NF (per)</td>
</tr>
<tr>
<td>Filler</td>
<td>CaC(¾) (per)</td>
</tr>
<tr>
<td>Plastisizer</td>
<td>Palatinol N (per)</td>
</tr>
<tr>
<td>Defoamer</td>
<td>BYK3155 (per)</td>
</tr>
<tr>
<td>Blowing agent</td>
<td>Func. Bicar. (per)</td>
</tr>
<tr>
<td>Gelation temp (°C)</td>
<td>200 210 210 200 200 210 210</td>
</tr>
<tr>
<td>Gelation Time (sec.)</td>
<td>90 90 120 90 120 90 120</td>
</tr>
<tr>
<td>Thickness (mm)</td>
<td>2.22 2.44 2.08 2.60 2.89 3.12 2.92</td>
</tr>
<tr>
<td>Expansion ratio (%)</td>
<td>300 330 280 350 390 420 390</td>
</tr>
<tr>
<td>Density (g/cm³)</td>
<td>0.51 0.45 0.53 0.42 0.38 0.35 0.39</td>
</tr>
</tbody>
</table>
Example 22 (not in accordance with invention)

Functionalization Additive = none

A commercial sodium bicarbonate (SB/03 from Solvay) was milled to reduce the particle size. The sample is identified as Example 22A (without functionalization additive).

For Example 22, PVC Plastisol compositions were prepared in a Dispermat apparatus. In each example, the components of the plastisols were dispersed on a high speed mixer and then mixed and de-aerated under vacuum. The resulting liquid PVC pastes (plastisols) are in each case spread coated at a thickness of about 0.75 mm on a piece of paper (Claire Fontaine) and heated in a ThermoSol Werner Mathis oven set at 200 or 210 °C for 90 or 120 seconds.

The expansion ratio was calculated as the ratio of the thickness of the foamed product after Werner Mathis oven heating over the initial thickness of the spread coated layer (here about 0.75 mm) before it passed the Werner Mathis oven heating. The specific components and amounts in each plastisol, the oven temperature, the time for gelation, the thickness of the foamed product after oven heating, the expansion ratio and the density of the PVC foams are provided in TABLE 11, where "per" means parts by weight of PVC resin.
TABLE 11: Without functionalization additive

<table>
<thead>
<tr>
<th>Example</th>
<th>22</th>
</tr>
</thead>
<tbody>
<tr>
<td>D10/D50/D90 (µm)</td>
<td>1.3/7/16</td>
</tr>
<tr>
<td>Plastisol Ex.</td>
<td>22 p1</td>
</tr>
<tr>
<td>PVC resin</td>
<td>367 NF (pcr)</td>
</tr>
<tr>
<td>Filler</td>
<td>CaCO₃ (pcr)</td>
</tr>
<tr>
<td>Plastisizer</td>
<td>Palatinol N (pcr)</td>
</tr>
<tr>
<td>Defoamer</td>
<td>BYK3155 (pcr)</td>
</tr>
<tr>
<td>Blowing agent</td>
<td>Non-Func. Bicar. (pcr)</td>
</tr>
<tr>
<td>Gelation temp (°C)</td>
<td>200</td>
</tr>
<tr>
<td>Gelation Time (sec.)</td>
<td>90</td>
</tr>
<tr>
<td>Thickness (mm)</td>
<td>1.22</td>
</tr>
<tr>
<td>Expansion ratio (%)</td>
<td>160</td>
</tr>
<tr>
<td>Density (g/cm³)</td>
<td>0.90</td>
</tr>
</tbody>
</table>

Examples 23-24 (in accordance with invention)

amorphous precipitated silica +

Functionalization Additive = PEG400

Two samples of functionalized sodium bicarbonate were prepared by extrusion of sodium bicarbonate particles (with a D90 = 20 microns) with PEG400 as the polyoxyalkylene additive and amorphous precipitated silica (Tixosil® 38AB by Rhodia). The functionalized sodium bicarbonate samples are identified as Example 23A (15 wt% of 93:7 PEG400 and Tixosil® 38AB) and Example 24A (30 wt% of 97:3 PEG400 and Tixosil® 38AB).

For Examples 24-25, PVC Plastisol compositions were prepared in a Dispermat apparatus. In each example, the components of the plastisols were dispersed on a high speed mixer and then mixed and de-aerated under vacuum. The resulting liquid PVC pastes (plastisols) of Example 8 are in each case spread coated at a thickness of about 0.75 mm on a piece of paper (SAPPI) and heated in a Thermosol Werner Mathis oven set at 200 or 210 °C for 90 or 120 seconds. The expansion ratio was calculated as the ratio of the thickness of the foamed product after Werner Mathis oven heating over the initial thickness of the spread coated layer (here about 0.75 mm) before it passed the Werner Mathis oven heating. The specific components and amounts in each plastisol, the oven temperature, the time for gelation, the thickness of the foamed product after oven heating, the expansion ratio and the
density of the PVC foams are provided in TABLE 12, where “per” means parts by weight of PVC resin.

**TABLE 12 : Functionalization Additive = PEG400 ; with amorphous precipitated silica**

<table>
<thead>
<tr>
<th>Example</th>
<th>23</th>
<th>24</th>
</tr>
</thead>
<tbody>
<tr>
<td>Functionalized Bicarbonate</td>
<td>Ex. 23A</td>
<td>Ex. 24A</td>
</tr>
<tr>
<td>Funct. Method</td>
<td>Extrusion</td>
<td>Extrusion</td>
</tr>
<tr>
<td>Funct. additive</td>
<td>13.95 wt% PEG400</td>
<td>29.1 wt% PEG400</td>
</tr>
<tr>
<td>Other component</td>
<td>1.05 wt% Tixosil® 38AB silica</td>
<td>0.9 wt% Tixosil® 38AB silica</td>
</tr>
<tr>
<td>Resulting Blowing agent Composition</td>
<td>85 wt% Na bicarbonate / 13.95 wt% PEG400 / 1.05 wt% Tixosil® 38 AB silica</td>
<td>70 wt% Na bicarbonate / 29.1 wt% PEG400 / 0.9 wt% Tixosil® 38 AB silica</td>
</tr>
<tr>
<td>Plastisol Ex.</td>
<td>23 p1</td>
<td>24 p1</td>
</tr>
<tr>
<td>PVC resin</td>
<td>367 NK (pcr)</td>
<td>100</td>
</tr>
<tr>
<td>Filler</td>
<td>CaCO₃ (pcr)</td>
<td>120</td>
</tr>
<tr>
<td>Plastisizer</td>
<td>Palatinol N (pcr)</td>
<td>80</td>
</tr>
<tr>
<td>Blowing agent</td>
<td>Func. Bicar. (pcr)</td>
<td>9.4</td>
</tr>
<tr>
<td>Gelation temp (°C)</td>
<td>200</td>
<td>200</td>
</tr>
<tr>
<td>Gelation Time (sec.)</td>
<td>90</td>
<td>90</td>
</tr>
<tr>
<td>Thickness (mm)</td>
<td>2.5</td>
<td>2.4</td>
</tr>
<tr>
<td>Expansion ratio (%)</td>
<td>330</td>
<td>310</td>
</tr>
<tr>
<td>Density (g/cm³)</td>
<td>0.4</td>
<td>0.4</td>
</tr>
</tbody>
</table>

Examples 25-26 (in accordance with invention)

**Functionalization Polyoxyalkylene Additive = Polypropylene Glycol (PPG400)**

Two samples of functionalized sodium bicarbonate were prepared by extrusion/milling of sodium bicarbonate particles with a polypropylene glycol of low molecular weight (<1000 g/mol): PPG400 as the functionalization additive. The functionalized sodium bicarbonate samples are identified as Example 25A (15 wt% PPG400) and Example 26A (30wt% PPG400).

For Examples 25-26, the same method of preparation of the PVC Plastisol compositions and preparation of PVC foams were used as described for Example 8 except that the oven for PVC foam production was an Hermatherm oven. The expansion ratio was calculated as the ratio of the thickness of the foamed product after the Hermatherm oven heating over the initial thickness of the spread coated layer (here about 0.75 mm) before it
passed the Hermatherm oven heating. The specific components and amounts in each plastisol, the oven temperature, the time for gelation, and the expansion ratio of the foamed PVC are provided in **TABLE 13**, where "per" means parts by weight of PVC resin.

### 5. **TABLE 13**: Functionalization Polyoxyalkylene Additive = Polypropylene Glycol (PPG400)

<table>
<thead>
<tr>
<th>Example</th>
<th>25</th>
</tr>
</thead>
<tbody>
<tr>
<td>Functionalized Bicarbonate</td>
<td>Ex. 25A</td>
</tr>
<tr>
<td>Funct. Method</td>
<td>Extrusion/milling</td>
</tr>
<tr>
<td>Funct. additive</td>
<td>15 wt% PPG400</td>
</tr>
<tr>
<td>Resulting Blowing agent Composition</td>
<td>85 wt% Na bicarbonate / 15 wt% PPG400</td>
</tr>
<tr>
<td>Plastisol Ex.</td>
<td>25 p1 25 p2 25 p3 25p4</td>
</tr>
<tr>
<td>PVC resin</td>
<td>367 NF (per) 100 100 100 100</td>
</tr>
<tr>
<td>Filler</td>
<td>CaCO₃ (per) 120 120 120 120</td>
</tr>
<tr>
<td>Plastisizer</td>
<td>Palatinol N (per) 80 80 80 80</td>
</tr>
<tr>
<td>Defoamer</td>
<td>BYK3155 (per) - - - -</td>
</tr>
<tr>
<td>Blowing agent</td>
<td></td>
</tr>
<tr>
<td>Func. Bicar. (per)</td>
<td>9.4 9.4 9.4 9.4</td>
</tr>
<tr>
<td>Incl. NaHCO₃ (per)</td>
<td>8 8 8 8</td>
</tr>
<tr>
<td>Gelation temp (°C)</td>
<td>200 200 210 210</td>
</tr>
<tr>
<td>Gelation Time (sec.)</td>
<td>90 120 90 120</td>
</tr>
<tr>
<td>Expansion ratio (%)</td>
<td>250 190 250 200</td>
</tr>
<tr>
<td>Density (g/cm³)</td>
<td>0.6 0.7 0.5 0.7</td>
</tr>
</tbody>
</table>

### 26

<table>
<thead>
<tr>
<th>Example</th>
<th>26</th>
</tr>
</thead>
<tbody>
<tr>
<td>Functionalized Bicarbonate</td>
<td>Ex. 26A</td>
</tr>
<tr>
<td>Funct. Method</td>
<td>Extrusion/milling</td>
</tr>
<tr>
<td>Funct. additive</td>
<td>30 wt% PPG400</td>
</tr>
<tr>
<td>Resulting Blowing agent Composition</td>
<td>70 wt% Na bicarbonate / 30 wt% PPG400</td>
</tr>
<tr>
<td>Plastisol Ex.</td>
<td>26 p1 26 p2 26 p3 26 p4</td>
</tr>
<tr>
<td>PVC resin</td>
<td>367 NF (per) 100 100 100 100</td>
</tr>
<tr>
<td>Filler</td>
<td>CaCC₃ (per) 120 120 120 120</td>
</tr>
<tr>
<td>Plastisizer</td>
<td>Palatinol N (per) 80 80 80 80</td>
</tr>
<tr>
<td>Defoamer</td>
<td>BYK3155 (per) - - - -</td>
</tr>
<tr>
<td>Blowing agent</td>
<td></td>
</tr>
<tr>
<td>Func. Bicar. (per)</td>
<td>11.4 11.4 11.4 11.4</td>
</tr>
<tr>
<td>Incl. NaHCO₃ (per)</td>
<td>8 8 8 8</td>
</tr>
<tr>
<td>Gelation temp (°C)</td>
<td>200 200 210 210</td>
</tr>
<tr>
<td>Gelation Time (sec.)</td>
<td>90 120 90 120</td>
</tr>
<tr>
<td>Expansion ratio (%)</td>
<td>270 240 250 180</td>
</tr>
<tr>
<td>Density (g/cm³)</td>
<td>0.5 0.5 0.5 0.7</td>
</tr>
</tbody>
</table>
Examples 27-28 (in accordance with invention)

Functionalization Polyoxyalkylene Additive = Polypropylene Glycol (PPG425)
with amorphous precipitated silica

Two samples of functionalized sodium bicarbonate were prepared by extrusion/milling of sodium bicarbonate particles with a polypropylene glycol of low molecular weight (<1000 g/mol): PPG425 as the functionalization additive and amorphous precipitated silica (Tixosil® 38AB by Rhodia). The functionalized sodium bicarbonate samples are identified as Example 27A (14wt% PPG425 and 1wt% Tixosil® 38AB) and Example 28A (29wt% PPG425 and 1wt% Tixosil® 38AB).

For Examples 27-28, the same method of preparation of the PVC Plastisol compositions and preparation of PVC foams were used as described for Example 8 except that the oven for PVC foam production was an Hermatherm oven. The expansion ratio was calculated as the ratio of the thickness of the foamed product after the Hermatherm oven heating over the initial thickness of the spread coated layer (here about 0.75 mm) before it passed the Hermatherm oven heating. The specific components and amounts in each plastisol, the oven temperature, the time for gelation, and the expansion ratio of the foamed PVC are provided in TABLE 14, where "per" means parts by weight of PVC resin.

TABLE 14: Functionalization Polyoxyalkylene Additive = Polypropylene Glycol (PPG425) ; with amorphous precipitated silica

<table>
<thead>
<tr>
<th>Example</th>
<th>27</th>
</tr>
</thead>
<tbody>
<tr>
<td>Functionalized Bicarbonate</td>
<td>Ex. 27A</td>
</tr>
<tr>
<td>Funct. Method</td>
<td>Extrusion/milling</td>
</tr>
<tr>
<td>Funct. additive</td>
<td>14 wt% PPG425</td>
</tr>
<tr>
<td>Other component</td>
<td>1 wt% Tixosil® 38AB silica</td>
</tr>
<tr>
<td>Resulting Blowing agent Composition</td>
<td>85 wt% Na bicarbonate / 14 wt% PPG425/ 1 wt% Tixosil® 38 AB silica</td>
</tr>
<tr>
<td>Plastisol Ex.</td>
<td>27</td>
</tr>
<tr>
<td>PVC resin</td>
<td>367 NF (per)</td>
</tr>
<tr>
<td>Filler</td>
<td>CaCC3 (per)</td>
</tr>
<tr>
<td>Plastisizer</td>
<td>Palatinol N (per)</td>
</tr>
<tr>
<td>Defoamer</td>
<td>BYK3155 (per)</td>
</tr>
<tr>
<td>Blowing agent</td>
<td>Func. Bicar. (per)</td>
</tr>
<tr>
<td></td>
<td>Incl. NaHCC (per)</td>
</tr>
<tr>
<td>Gelation temp (°C)</td>
<td>200</td>
</tr>
<tr>
<td>Gelation Time (sec.)</td>
<td>90</td>
</tr>
<tr>
<td>Expansion ratio (%)</td>
<td>280</td>
</tr>
<tr>
<td>---------------------</td>
<td>-----</td>
</tr>
<tr>
<td>Density (g/cm³)</td>
<td>0.4</td>
</tr>
</tbody>
</table>

**Example 28**

<table>
<thead>
<tr>
<th>Functionalized Bicarbonate</th>
<th>Ex. 28A</th>
</tr>
</thead>
<tbody>
<tr>
<td>Func. Method</td>
<td>Extrusion/milling</td>
</tr>
<tr>
<td>Func. additive</td>
<td>15 wt% PPG425</td>
</tr>
<tr>
<td>Other component</td>
<td>1 wt% Tixosil® 38AB silica</td>
</tr>
<tr>
<td>Resulting Blowing agent Composition</td>
<td>70 wt% Na bicarbonate / 29 wt% PPG425 / 1 wt% Tixosil® 38 AB silica</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Plastisol Ex.</th>
<th>28 p1</th>
<th>28 p2</th>
<th>28 p3</th>
<th>28 p4</th>
</tr>
</thead>
<tbody>
<tr>
<td>PVC resin</td>
<td>367 NF (per)</td>
<td>100</td>
<td>100</td>
<td>100</td>
</tr>
<tr>
<td>Filler</td>
<td>CaCO₃ (per)</td>
<td>120</td>
<td>120</td>
<td>120</td>
</tr>
<tr>
<td>Plastisizer</td>
<td>Palatinol N (per)</td>
<td>80</td>
<td>80</td>
<td>80</td>
</tr>
<tr>
<td>Defoamer</td>
<td>BYK3155 (per)</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Blowing agent</td>
<td>Func. Bicar. (per)</td>
<td>11.4</td>
<td>11.4</td>
<td>11.4</td>
</tr>
<tr>
<td></td>
<td>Incl. NaHCO₃ (per)</td>
<td>8</td>
<td>8</td>
<td>8</td>
</tr>
<tr>
<td>Gelation temp (°C)</td>
<td>200</td>
<td>200</td>
<td>210</td>
<td>210</td>
</tr>
<tr>
<td>Gelation Time (sec.)</td>
<td>90</td>
<td>120</td>
<td>90</td>
<td>120</td>
</tr>
<tr>
<td>Expansion ratio (%)</td>
<td>330</td>
<td>300</td>
<td>310</td>
<td>220</td>
</tr>
<tr>
<td>Density (g/cm³)</td>
<td>0.4</td>
<td>0.4</td>
<td>0.4</td>
<td>0.6</td>
</tr>
</tbody>
</table>

The disclosure of all patent applications, and publications cited herein are hereby incorporated by reference, to the extent that they provide exemplary, procedural or other details supplementary to those set forth herein.

Should the disclosure of any of the patents, patent applications, and publications that are incorporated herein by reference conflict with the present specification to the extent that it might render a term unclear, the present specification shall take precedence.

In the present application, where an element or component is said to be included in and/or selected from a list of recited elements or components, it should be understood that in related embodiments explicitly contemplated here, the element or component can also be any one of the individual recited elements or components, or can also be selected from a group consisting of any two or more of the explicitly listed elements or components. Any element or component recited in a list of elements or components may be omitted from such list. Further, it should be understood that elements, embodiments, and/or features of processes or
methods described herein can be combined in a variety of ways without departing from the scope and disclosure of the present teaching, whether explicit or implicit herein.

Accordingly, the scope of protection is not limited by the description set out above, but is only limited by the claims which follow, that scope including all equivalents of the subject matter of the claims. Each and every claim is incorporated into the specification as an embodiment of the present invention. Thus, the claims are a further description and are an addition to the preferred embodiments of the present invention.

While preferred embodiments of this invention have been shown and described, modifications thereof can be made by one skilled in the art without departing from the spirit or teaching of this invention. The embodiments described herein are exemplary only and are not limiting. Many variations and modifications of systems and methods are possible and are within the scope of the invention.

We claim:
CLAIMS

1. A chemical blowing agent for foaming a thermoplastic polymer precursor, for example PVC plastisol or a polymer resin in an extrusion process, said chemical blowing agent comprising a functionalized particulate bicarbonate, wherein said functionalized particulate bicarbonate contains at least one functionalization additive, and wherein said functionalization additive in the functionalized particulate bicarbonate comprises a polyoxyalkylene additive selected from the group consisting of polyoxyalkylenes, any derivatives thereof (such as esters), salts thereof, and any combination of two or more thereof.

2. The chemical blowing agent according to Claim 1, wherein the blowing agent does not contain any further blowing agent which is an exothermic blowing agent, or wherein the chemical blowing agent does not contain a compound which liberates nitrogen gas during heating, or wherein the chemical blowing agent does not contain a compound which liberates ammonia gas during heating.

3. The chemical blowing agent according to Claim 1 or 2, wherein the polyoxyalkylene additive comprises:
   a polyethylene oxide (PEO) polymer, preferably of a MW of 20,000 or more,
   a polyethylene glycol (PEG) polymer, preferably of a MW of less than 20,000, or a MW of 4000 or less, or a MW of 3600 or less, or a MW of 2000 or less, or a MW of 1000 or less, or a MW of 800 or less,
   a polypropylene oxide (PPO) polymer,
   a polypropylene glycol (PPG) polymer,
   a co-polymer or block polymer which is made with at least one monomer of propylene oxide or propylene glycol,
   a co-polymer or block polymer which is made with at least one monomer of ethylene oxide or ethylene glycol,
   a polyethylene glycol / polypropylene glycol (PEG/PPG) block co-polymer, such as the PEG/PPG/PEG triblock polymer, or any derivatives (such as esters) thereof,
wherein any of these polymers may be chemically functionalized or not.

4. The chemical blowing agent according to any of Claims 1 to 3, wherein the polyoxyalkylene additive comprises a polyethylene glycol, preferably a polyethylene glycol with a molecular weight less than 1000 g/mol.

5. The chemical blowing agent according to any of Claims 1 to 3, wherein the polyoxyalkylene additive comprises a polyoxyalkylene derivative which is an ester, such as a PEG or PPG ester.

6. The chemical blowing agent according to any of Claims 1 to 3, wherein the polyoxyalkylene additive is a polyoxyalkylene derivative which is a chemically functionalized polyoxyalkylene, for example, a polyoxyalkylene bearing one or more reactive substituents, such as -OH, -NH₂, -NHR, -SH and -COOH, wherein R is a hydrocarbon residue, and which has been reacted with a functional chemical compound being reactive with the reactive substituent of the polyoxyalkylene.

7. The chemical blowing agent according to any of Claims 1 to 6, wherein the functionalized particulate bicarbonate comprises at least one additional functionalization additive selected from
- one or more polymers other than the polyoxyalkylene additive;
- one or more amino acids, any derivative thereof, and salts thereof;
- one or more inorganic salts;
- one or more oils;
- one or more fats;
- one or more resin acids, any derivative thereof, and salts thereof;
- one or more fatty acids, any derivative thereof, and salts thereof;
- a carboxylic or polycarboxylic acid, derivative thereof (such as esters), or salts thereof;
- one or more soaps;
- one or more waxes; or
- any combinations thereof;

preferably selected from the group consisting of:
- one or more fatty acids, any derivative thereof, and salts thereof;
8. The chemical blowing agent according to any of Claims 1 to 7, wherein the functionalized particulate bicarbonate comprises at least 50% by weight and less than 100% by weight of the bicarbonate component, and from 50% or less to 0.02% by weight of at least one of said functionalization additive, preferably comprises at least 65% by weight and less than 100% by weight of the bicarbonate component, and from 35% or less to 0.02% by weight of at least one of said functionalization additive.

9. The chemical blowing agent according to any of Claims 1 to 8, wherein the particles of the functionalized particulate bicarbonate have a particle size distribution of D_{50} of more than 1 µm and at most 250 µm, preferably at most 100 µm, more preferably at most 60 µm, yet more preferably at most 40 µm, yet most preferably at most 25 µm.

10. The chemical blowing agent according to any of Claims 1 to 9, wherein the functionalized particulate bicarbonate is obtained by at least one of the following processes:
   - by spray-drying (also known as atomization), wherein the additive is dissolved in the bicarbonate-containing solution.
   - by grinding or co-grinding (also known as milling or co-milling) with the additive(s) in emulsion or powder form;
   - by spray coating and granulation within a fluidized bed,
   - by spray agglomeration within a fluidized bed,
   - by spray chilling (e.g., spray cooling, spray freezing),
   - by roller compaction, and/or
   - by extrusion, including simultaneous mixing/extrusion:
     optionally followed by subjecting the thus obtained functionalized particulate bicarbonate to milling to reduce its mean particle size.

11. The chemical blowing agent according to any of Claims 1 to 9, further comprising an second compound which liberates C02 upon heating, said second compound being selected from the group consisting of a carboxylic or polycarboxylic acid, derivative thereof (such as esters), or salts thereof, wherein said second compound optionally is functionalized with at
least one additive which is different or the same as the one in the functionalized particulate bicarbonate, preferably the same additive, and wherein said second compound preferably is at least one of:
- fumaric acid,
- tartaric acid,
- citric acid, citrates (such as sodium hydrogen citrate, disodium citrate), or esters of citric acid; or combination thereof.

12. The chemical blowing agent according to any of Claims 1 to 11, wherein the functionalized particulate bicarbonate further comprises silica, preferably comprises amorphous silica, more preferably comprises amorphous precipitated silica.

13. A foamable polymer composition, comprising a polymer, optionally a foaming stabilizer, and the chemical blowing agent according to any of Claims 1 to 12, wherein the polymer is a PVC, a polyurethane, a polyolefin, or a polyamide.

14. The foamable polymer composition according to Claim 13, further comprising an exothermic chemical blowing agent, preferably selected from the group consisting of ADCA (azodicarbonamide), OBSH (4,4’-oxybis(benzenesulfonyl hydrazide)), DNPT (dinitrosopentamethylene tetramine), PTSS (p-toluene semicarbazide), BSH (benzene-4-sulfonohydrazide), and TSH (toluene-4-sulfonohydrazide), 5-PT (5-phenyltetrazole), more preferably comprising ADCA.

15. The foamable polymer composition according to Claim 12 or 14, being a foamable PVC plastisol composition, wherein the polymer comprises a PVC resin.

16. The foamable polymer composition according to Claim 12, being a foamable PVC plastisol composition, wherein the polymer comprises a PVC resin, which preferably does not contain azodicarbonamide, benzenesulfonylhydrazide, and/or p-toluenesulfonilhydrazide.

17. A process for manufacturing a polymer, comprising heating the foamable polymer composition according to Claim 13 or 14 which comprises the functionalized particulate bicarbonate at a temperature suitable for liberating CO2 gas and melting the polymer during
a gelation time at a temperature above glass transition temperature $T_g$ and/or above melting temperature $T_m$ of the polymer which is less than 130 seconds.

18. A process for manufacturing a PVC polymer, comprising heating the foamable PVC polymer composition according to Claim 14 or 15 which comprises the functionalized particulate bicarbonate and the PVC resin at a temperature suitable for liberating CO₂ gas and melting the PVC polymer during a gelation time from 90 seconds to 120 seconds at a temperature from 190 to 210°C, preferably from 200 to 210°C, to provide a foamed PVC polymer, wherein the foamable polymer composition preferably is spread coated on a surface before heating and melting, and wherein the foamed polymer has an expansion ratio of at least 270, preferably at least 280, more preferably at least 300 and/or has a density of less than 0.6 g/cm³, preferably less than 0.55 g/cm³, more preferably at most 0.5 g/cm³.

19. A process for manufacturing a polymer, comprising heating the foamable PVC plastisol composition according to Claim 15 or 16 which comprises the functionalized particulate bicarbonate at a temperature suitable for liberating CO₂ gas and melting the PVC resin during a gelation time, wherein said gelation time with the chemical blowing agent comprising the functionalized particulate is less than the gelation time that would be obtained with a chemical blowing agent liberating N₂, all other components in the foamable PVC plastisol composition being the same.
INTERNATIONAL SEARCH REPORT

A. CLASSIFICATION OF SUBJECT MATTER
INV. C08J9/08 C08K3/26 C08K3/34
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)
C08J C08K

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, WPI Data

C. DOCUMENTS CONSIDERED TO BE RELEVANT

<table>
<thead>
<tr>
<th>Category*</th>
<th>Citation of document, with indication, where appropriate, of the relevant passages</th>
<th>Relevant to claim No.</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>col umn 2, line 20 - line 51</td>
<td>9, 12, 14-19</td>
</tr>
<tr>
<td></td>
<td>col umn 5, line 44 - line 45</td>
<td></td>
</tr>
<tr>
<td>X</td>
<td>WO 2009/038783 AI (SURMODICS INC [US]; REED PAMELA J [US]; ROLFES EMILY R [US])</td>
<td>1-8, 10</td>
</tr>
<tr>
<td></td>
<td>26 March 2009 (2009-03-26) page 11, line 5 - line 7</td>
<td>11</td>
</tr>
<tr>
<td></td>
<td>page 30, line 24 - line 27</td>
<td></td>
</tr>
<tr>
<td></td>
<td>page 43, line 18 - line 22</td>
<td></td>
</tr>
<tr>
<td></td>
<td>page 51, line 21 - page 52, line 17</td>
<td></td>
</tr>
<tr>
<td>Y</td>
<td>EP 2 508 555 AI (ELECTROLUX HOME PROD CORP [BE]) 10 October 2012 (2012-10-10)</td>
<td>12, 14, 15</td>
</tr>
<tr>
<td></td>
<td>paragraphs [0035], [0037], claim 4</td>
<td></td>
</tr>
</tbody>
</table>

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 document 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
A: document member of the same patent family

Date of the actual completion of the international search 22 August 2018

Date of mailing of the international search report 30/08/2018

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
Magri zo, Simeon

Form PCT/ISA/210 (second sheet) (April 2005)
<table>
<thead>
<tr>
<th>Category</th>
<th>Citation of document, with indication, where appropriate, of the relevant passages</th>
<th>Relevant to claim No.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Y</td>
<td>US 4 892 854 A (PASTORINO RONALD L [US] ET AL) 9 January 1990 (1990-01-09) claim 10; table 3</td>
<td>17, 18</td>
</tr>
<tr>
<td>Y</td>
<td>wo 90/14384 AI (HUBER CORP J M [US]) 29 November 1990 (1990-11-29) page 4, line 11 - line 22 page 7, line 8 - line 12 page 8, line 18 - line 21 page 9, line 1 - line 15 page 12, line 23 - line 33 page 13, line 8 - line 16 page 14, line 15 - line 18 page 25; table 1</td>
<td>16, 19</td>
</tr>
<tr>
<td>Y</td>
<td>EP 3 037 388 AI (SOLVAY [BE]) 29 June 2016 (2016-06-29) cited in the application paragraphs [0027], [0049], [0052], [0058] - [0062]</td>
<td>9</td>
</tr>
</tbody>
</table>

Form PCT/ISA/210 (continuation of second sheet) (April 2008)
<table>
<thead>
<tr>
<th>Patent document cited in search report</th>
<th>Publication date</th>
<th>Patent family member(s)</th>
<th>Publication date</th>
</tr>
</thead>
<tbody>
<tr>
<td>US 5047440</td>
<td>10-09-1991</td>
<td>NONE</td>
<td></td>
</tr>
<tr>
<td>WO 2009038783</td>
<td>26-03-2009</td>
<td>CA 2699868 A1</td>
<td>26-03-2009</td>
</tr>
<tr>
<td></td>
<td></td>
<td>EP 2185637 A1</td>
<td>19-05-2010</td>
</tr>
<tr>
<td></td>
<td></td>
<td>JP 5474798 B2</td>
<td>16-04-2014</td>
</tr>
<tr>
<td></td>
<td></td>
<td>JP 2010538791 A</td>
<td>16-12-2010</td>
</tr>
<tr>
<td></td>
<td></td>
<td>US 2009093550 A1</td>
<td>09-04-2009</td>
</tr>
<tr>
<td></td>
<td></td>
<td>WO 2009038783 A1</td>
<td>26-03-2009</td>
</tr>
<tr>
<td>EP 2508555</td>
<td>10-10-2012</td>
<td>NONE</td>
<td></td>
</tr>
<tr>
<td>US 4892854</td>
<td>09-01-1990</td>
<td>DE 69017722 D1</td>
<td>20-04-1995</td>
</tr>
<tr>
<td></td>
<td></td>
<td>DE 69017722 T2</td>
<td>14-09-1995</td>
</tr>
<tr>
<td></td>
<td></td>
<td>EP 0377507 A2</td>
<td>11-07-1990</td>
</tr>
<tr>
<td></td>
<td></td>
<td>JP 3088008 B2</td>
<td>18-09-2000</td>
</tr>
<tr>
<td></td>
<td></td>
<td>JP H0328235 A</td>
<td>06-02-1991</td>
</tr>
<tr>
<td></td>
<td></td>
<td>US 4892854 A</td>
<td>09-01-1990</td>
</tr>
<tr>
<td>WO 9014384</td>
<td>29-11-1990</td>
<td>AT 130331 T</td>
<td>15-12-1995</td>
</tr>
<tr>
<td></td>
<td></td>
<td>AU 635035 B2</td>
<td>11-03-1993</td>
</tr>
<tr>
<td></td>
<td></td>
<td>CA 2016822 A1</td>
<td>16-11-1990</td>
</tr>
<tr>
<td></td>
<td></td>
<td>DE 69023649 D1</td>
<td>21-12-1995</td>
</tr>
<tr>
<td></td>
<td></td>
<td>JP H04506677 A</td>
<td>19-11-1992</td>
</tr>
<tr>
<td></td>
<td></td>
<td>NO 914494 A</td>
<td>10-01-1992</td>
</tr>
<tr>
<td></td>
<td></td>
<td>WO 9014384 A1</td>
<td>29-11-1990</td>
</tr>
<tr>
<td></td>
<td></td>
<td>BR 112017013484 A2</td>
<td>27-02-2018</td>
</tr>
<tr>
<td></td>
<td></td>
<td>CA 2970111 A1</td>
<td>30-06-2016</td>
</tr>
<tr>
<td></td>
<td></td>
<td>CN 107108246 A</td>
<td>29-08-2017</td>
</tr>
<tr>
<td></td>
<td></td>
<td>EP 3037388 A1</td>
<td>29-06-2016</td>
</tr>
<tr>
<td></td>
<td></td>
<td>EP 3237330 A1</td>
<td>01-11-2017</td>
</tr>
<tr>
<td></td>
<td></td>
<td>JP 2018502038 A</td>
<td>25-01-2018</td>
</tr>
<tr>
<td></td>
<td></td>
<td>KR 20170100554 A</td>
<td>04-09-2017</td>
</tr>
<tr>
<td></td>
<td></td>
<td>US 2017349446 A</td>
<td>07-12-2017</td>
</tr>
<tr>
<td></td>
<td></td>
<td>WO 2016102603 A1</td>
<td>30-06-2016</td>
</tr>
</tbody>
</table>