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SIOC BARRIER LAYER TO ALKALI METALS AND PROCESS OF MAKING BARRIER LAYER TO SIOC ALKALI METALS

The present invention relates to a thin silicon-oxycarbide-based transparent layer that forms a barrier to the diffusion of alkali-metal ions. The invention also relates to a process for manufacturing such a layer on a mineral glass substrate.

It is known that the diffusion of alkali-metal ions, in particular of sodium ions, from substrates made of glass (borosilsicate glass, soda-lime glass) toward and into thin functional layers such as TiO₂-based photocatalytic layers, transparent conductive oxide (TCO) layers or hydrophobic organic layers deteriorates the advantageous properties inherent in these functional layers.

It is also known to interpose, between the substrate made of mineral glass rich in alkalimetal ions and said functional layer to be protected, a thin layer of a mineral oxide as a barrier against the diffusion of alkali-metal ions. Mention may be made, as examples of such oxides, of silica, alumina, zirconia, zinc/tin oxide, titanium oxide and silicon oxycarbide (SiOC). The barrier properties to alkali-metal ions of silicon oxycarbide layers are mentioned, for example, in applications EP 1 708 812, EP 1 686 595, EP 275 662 and US 2003/ 612033.

To the Applicant's knowledge, no study exists at the current time regarding the influence of the carbon content of SiOC layers on the ability to block or capture alkali-metal ions. In the context of its studies that aim to optimize silicon oxycarbide sublayers with a view to their subsequent functionalization by hydrophobic agents, the Applicant observed that the silicon oxycarbide layers richest in carbon had the best barrier properties to alkalimetal ions and were also excellent tie layers for the hydrophobic functional agents.

Increasing the carbon content of the thin silicon oxycarbide layers resulted, however, in the undesirable appearance of a yellow coloration of the glazings. Such a coloration is problematic from an esthetic viewpoint, in particular in the field of motor

vehicle glazings and glazings for dwellings. It is an even greater problem in the field of photovoltaic cells, where it significantly reduces energy efficiency by filtering out the blue and green components of visible light.

The present invention is based on the discovery that it was possible to significantly reduce the yellow color of thin carbon-rich silicon oxycarbide layers by depositing directly onto these layers a silicon oxycarbide layer significantly poorer in carbon than the underlying layer.

The present invention therefore relates to a glass substrate containing alkali-metal ions and provided with a first transparent thin carbon-rich silicon oxycarbide layer and with a

second transparent thin layer, deposited directly onto the first layer and which is significantly less rich in carbon than the first layer.

As it is, however, often impossible to determine, in the final product, the boundary between the carbon-rich SiOC layer and the carbon-poor or carbon-free layer, these two layers are defined in the present application as a single layer with a carbon-rich deep zone (corresponding to the first layer mentioned above) and a carbon-poor surface zone (corresponding to the second layer mentioned above).

One subject of the present invention is consequently a glazing comprising a transparent glass substrate containing ions of at least one alkali metal and a transparent layer made of silicon oxycarbide (SiO_xC_y) having a total thickness E with

- a carbon-rich deep zone, extending from a depth P_3 to a depth P_4 , where the C/Si atomic ratio is greater than or equal to 0.5, and
- a carbon-poor surface zone, extending from a depth P_1 to a depth P_2 , where the C/Si atomic ratio is less than or equal to 0.4,

with $P_1 < P_2 < P_3 < P_4$ and $(P_2-P_1) + (P_4-P_3) < E$

the distance between P_1 and P_2 representing from 10% to 70%, preferably from 15% to 50% and in particular from 25% to 40%, of the total thickness E of the silicon oxycarbide layer $(0.1 \le (P_2-P_1)/E \le 0.7)$ and

the distance between P_3 and P_4 representing from 10% to 70%, preferably from 25% to 60%, in particular from 35% to 50%, of the total thickness E of the silicon oxycarbide layer $(0.1 \le (P_4 - P_3)/E \le 0.7)$.

The sum of the distance between P_1 and P_2 and the distance between P_3 and P_4 is of course always less than the overall thickness E of the silicon oxycarbide layer since there exists, between P_2 and P_3 , a zone where the C/Si ratio is intermediate between 0.4 and 0.5, and, beyond P_4 , a zone where the C/Si ratio is less than 0.5. Furthermore, the zone nearest the surface, located at a depth of less than P_1 , generally has an extremely high C/Si ratio due to the presence of CO_2 originating from the atmosphere or from organic pollutants absorbed at the surface.

The silicon oxycarbide layer of the glazing of the present invention is a dense non-porous layer having a refractive index between 1.45 and 1.9.

Its total thickness E is preferably between 10 and 200 nm, more preferably between 20 and 100 nm, and in particular between 40 and 70 nm.

The absolute methods that make it possible to determine the value of the total thickness of the silicon oxycarbide layer are X-ray reflectometry and transmission electron microscopy (TEM).

In the appended figures, this total thickness of the SiO_xC_y layer is systematically indicated by an arrow. In these figures, and in particular with a view to determining the ratios $(P_2-P_1)/E$ and $(P_4-P_3)/E$, it is determined not by one of the absolute methods mentioned above, but by secondary ionization mass spectroscopy (SIMS), which makes it possible to obtain the variation in the silica (SiO_2) content as a function of the analysis time and therefore of the depth of the sample analyzed.

In the present invention, the interface between the substrate and the silicon oxycarbide layer is defined as being the depth where the concentration of silica is exactly intermediate between that of the glass substrate and that at the local minimum in the vicinity of the interface. The determination of this depth, corresponding to the total thickness E of the layer, is illustrated in figure 2, which shows the variation in the silica (SiO₂) content of the sample from figure 1. The silica content of the mineral glass substrate is equal to SiO_{2S} and that corresponding to the local minimum in the vicinity of the interface is equal to SiO_{2M}. The interface between the glass substrate and the layer deposited by CVD is defined as being located at the depth where the silica content (SiO_{2intermediate}) is equidistant between SiO_{2S} and SiO_{2M}.

This depth E corresponding to SiO_{2intermediate} has been reported on figure 1.

The carbon content of the silicon oxycarbide layer is also determined by SIMS. It is expressed in the present application by means of the ratio of the number of carbon atoms to the number of silicon atoms (C/Si atomic ratio).

Figure 1 shows, by way of example, the C/Si ratio as a function of the abrasion time (during the SIMS analysis) of a sample according to the invention. As is known, the abrasion time (indicated on the x axis) is proportional to the depth analyzed, the proportionality factor depending on the experimental conditions and on the nature and properties of the sample. It is observed on this figure that the C/Si ratio is very high at the surface due to possible carbon-containing pollutants and/or adsorbed CO_2 originating from the atmosphere. In the layers near the surface, between P_1 and P_2 , it is relatively low (of the order of 0.2) and then increases with the depth to exceed the value of 0.4 at a depth P_2 , and the value of 0.5 starting from a depth P_3 . It remains above this value until a depth P_4 and then tends toward zero upon arriving at the glass substrate.

As already explained in the introduction, the present invention is based on the discovery that the yellow color of thin carbon-rich silicon oxycarbide layers may be attenuated, or even eliminated, by depositing a thin carbon-poor silicon oxycarbide layer.

In the present application, this yellow color is quantified, as is known, using the CIE L*A*b* (hereinafter CIELab) colorimetric system from measurements made on the glazing, after deposition of the layers. The CIELab system defines a color space in the form of a sphere with an axis L* characterizing the lightness, a red/green axis a* and a blue/yellow axis b*. An a* value of greater than 0 corresponds to colors with a red component, a negative a* value to colors with a green component, a positive b* value to colors with a yellow component and a negative b* value to colors with a blue component.

Generally, gray-colored or colorless layers that have low a* and b* values, that is to say values close to zero, will be sought.

The glazings of the present invention preferably have a b* value (CIELab, illuminant D65) between -2 and +3, in particular between -1.5 and +2.0; and an a* value (CIEBab, illuminant D65) between -2 and +2, in particular between -1.5 and +1.5. These values apply to glazings with layers deposited on a transparent colorless substrate made of clear float glass sold by the company Saint-Gobain Glass France under the trade name SGG Planilux®.

Another subject of the present invention is a process for manufacturing the glazings described above.

This process essentially comprises the following two successive steps:

- (a) a first step of chemical vapor deposition (CVD) of a carbon-rich silicon oxycarbide layer onto at least one part of the surface of a mineral glass substrate, by bringing said surface into contact with a gas flow containing ethylene (C_2H_4), silane (SiH_4), carbon dioxide (CO_2) and nitrogen (N_2), at a temperature between $600^{\circ}C$ and $680^{\circ}C$, the ethylene/silane (C_2H_4/SiH_4) volume ratio during step (a) being less than or equal to 3.3,
- (b) a second step of chemical vapor deposition (CVD) of a carbon-poor silicon oxycarbide layer onto the carbon-rich layer obtained in step (a), with a gas flow containing ethylene (C₂H₄), silane (SiH₄), carbon dioxide (CO₂) and nitrogen (N₂), at a temperature between 600°C and 680°C, the ethylene/silane (C₂H₄/SiH₄) volume ratio during step (b) being less than or equal to 3.4.

The first step (step (a)) is preferably carried out on float glass inside the furnace on a tin bath or immediately after exit from the furnace. One particularly advantageous embodiment consists in carrying out the deposition by CVD inside the furnace, where it is relatively easy to create a confined atmosphere essential to a satisfactory control of the chemical composition of the reactive mixture and of the layer deposited.

The deposition temperatures indicated above are the temperatures of the tin bath measured in the immediate vicinity (less than 20 cm away) of the nozzle through which the reactive gases arrive.

When the two layers (steps (a) and (b)) are deposited by CVD, two nozzles are placed parallel to one another, at a distance generally between 2 and 5 m, the one upstream in the direction of travel of the ribbon forming the deepest layer/zone and the one downstream forming the least deep layer/zone.

Each of the nozzles preferably covers the entire width of the glass ribbon.

It is important to note that, although the process comprises two steps of forming a silicon oxycarbide layer, these "layers" are impossible to distinguish from one another in the final product, where they correspond to carbon-rich and carbon-poor zones of the single silicon oxycarbide layer.

The deposition of thin silicon oxycarbide layers by CVD is known and a person skilled in the art will know how to adjust the flow rates of the reactive gases as a function of the speed of travel of the glass, the furnace temperatures, and the thickness of the layers that it is desired to obtain. In the present invention, the experimental conditions of step (a) and optionally (b) are preferably adjusted such that the thickness of the SiO_xC_y layer deposited is, after step (b), between 10 and 200 nm, in particular between 20 and 100 nm and particularly preferably between 40 and 70 nm. These total thickness ranges are also preferred when a layer of silica is deposited below the carbon-rich SiO_xC_y zone by a method other than CVD.

The ethylene/silane (C_2H_4/SiH_4) volume ratio during step (a) is preferably between 1 and 3.3, in particular between 1.5 and 3.3. The ratio of oxidizing gas (CO_2) to silane (SiH_4) is generally between 1 and 50, preferably between 1.5 and 10 and in particular between 2 and 6.

For the deposition of the carbon-poor silicon oxycarbide layer (step (b)), the ethylene/silane (C_2H_4/SiH_4) volume ratio is preferably between 3.5 and 6, in particular between 3.7 and 5.5.

The process for manufacturing the glazings of the present invention preferably also comprises, after step (b), a third step (step (c)) of annealing and/or forming the

substrate bearing the silicon oxycarbide layer. During this step (c), the substrate is brought to a temperature between 580°C and 700°C, preferably between 600°C and 680°C.

Examples

A glazing according to the invention is prepared by CVD (C₂H₂, SiH₄, CO₂, N₂) with two parallel CVD nozzles having a width of 3.3 m each, onto a substrate made of clear float glass (Planilux®) respectively at a temperature of 648°C and 638°C (temperature of the tin bath in the immediate vicinity of the nozzle); the width of the ribbon is 3.6 m, its thickness is 2.5 mm and its speed of travel is 15 m/min. The two nozzles are inside the glass-making furnace, at a distance of 3 m from one another.

The first, upstream nozzle delivers the following mixture of reactive gases:

SiH₄: 14 nL/min, CO₂: 30 nL/min, C₂H₄: 39 nL/min (in diluent N₂) (C₂H₄/SiH₄ volume ratio = 2.79).

The second nozzle, downstream of the first, delivers the following mixture of reactive gases:

SiH₄: 14 nL/min, CO₂: 30 nL/min, C₂H₄: 52 nL/min (in diluent N₂) (C₂H₄/SiH₄ volume ratio = 3.71).

Comparative glazing is manufactured under the same experimental conditions. This comparative example differs from the example according to the invention simply by the fact that the second nozzle delivers a mixture of gases identical to that of the second nozzle, namely the following mixture:

SiH₄: 14 nL/min, CO₂: 30 nL/min, C₂H₄: 39 nL/min (in dilent N₂) (C₂H₄/SiH₄ volume ratio = 2.79).

All of the glazings are subjected to an annealing step at a temperature of 640°C for 8 minutes.

Figure 1 shows the SIMS spectrum of the glazing according to the invention where the $(P_2-P_1)/E$ ratio is equal to around 29% and the $(P_4-P_3)/E$ ratio is equal to around 46%.

Figure 3 shows the SIMS spectrum of the comparative example. The silicon oxycarbide layer of this sample has a high carbon content (C/Si > 0.4) both in the surface zone and in the deeper zone. It is impossible to determine the values of P_1 , P_2 and P_3 .

Table 1 presents the values of the parameters a* and b* (CIELab) for the glazings obtained according to the example and the comparative example.

It may be observed that the sample of the comparative example lacking a carbon-free or carbon-poor surface layer has a quite significant yellow color ($b^* = 3.52$). This value is

significantly reduced for the example according to the invention ($b^* = 1.39$) owing to the presence of the carbon-poor layer of the example according to the invention.

Table 1

	Example according to	Comparative
	the invention	example
Light transmission	82.08%	78.57%
a*	-0.95	-1.09
b*	1.39	3.52
Total thickness	60.8 nm	54 nm
determined by TEM		

BARRIERELAG AF SIOC-HOLDIGE ALKALIMETALLER PATENTKRAV

- 1. Rude, der omfatter et substrat af transparent glas, der indeholder ioner af mindst ét alkalimetal, og et transparent siliciumoxycarbidlag (SiO_xC_v), med en samlet tykkelse E med
 - (a) en dyb, carbonrig zone, der strækker sig fra en dybde P₃ til en dybde P₄, hvor det atomare forhold C/Si er større end eller lig med 0,5, og
 - (b) en carbonfattig overfladezone, der strækker sig fra en dybde P_1 til en dybde P_2 , hvor det atomare forhold C/Si er mindre end eller lig med 0,4,

med $P_1 < P_2 < P_3 < P_4$ og $(P_2-P_1) + (P_4-P_3) < E$,

hvor afstanden mellem P₁ og P₂ udgør fra 10 til 70 % af siliciumoxycarbidlagets samlede tykkelse E, og

afstanden mellem P₃ og P₄ udgør 10 til 70 % af siliciumoxycarbidlagets samlede tykkelse E.

- 2. Rude ifølge krav 1, kendetegnet ved, at SiO_xC-lagets samlede tykkelse ligger mellem 10 og 200 nm, fortrinsvis mellem 20 og 100 nm, især mellem 40 og 70 nm.
- 3. Rude ifølge krav 1 eller 2, kendetegnet ved, at det transparente siliciumoxycarbidlag er et ikke-porøst lag, der har et brydningsindeks mellem 1,45 et 1,9.
- 4. Rude ifølge et hvilket som helst af de foregående krav, kendetegnet ved, at den præsenterer en (CIELab)-værdi b* mellem -2 og +3, især mellem -1,5 og +2,0.
- 5. Fremgangsmåde til fremstilling af en rude ifølge et hvilket som helst af de foregående krav, kendetegnet ved, at den omfatter:
- (a) et første trin med kemisk dampudfældning (CVD) af et carbonrigt siliciumoxycarbidlag på mindst en firdel af overfladen af et mineralglassubstrat ved at bringe denne overflade i kontakt med en gasstrøm, der indeholder ethylen (C_2H_4), silan (SiH₄), carbondioxid (CO_2) og nitrogen (N_2), ved en temperatur mellem 600 °C og 680 °C, hvor volumenforholdet ethylen/silan (C_2H_4 /SiH₄) under trin (a) er mindre end eller lig med 3,3,
- (b) et andet trin med CVD af et carbonfattigt siliciumoxycarbidlag, på det carbonrige lag og opnået i trin (a), med en gasstrøm, der indeholder ethylen (C_2H_4), silan (SiH_4), carbondioxid (CO_2) og nitrogen (N_2), ved en temperatur mellem 600 °C og 680 °C, hvor volumenforholdet ethylen/silan (C_2H_4 / SiH_4) under trin (b) er større end 3,4.
- 6. Fremgangsmåde ifølge krav 5, kendetegnet ved, at den desuden omfatter et afhærdningstrin og/eller formningstrin ved en temperatur mellem 580 °C og 700 °C, fortrinsvis mellem 600 °C og 680 °C.

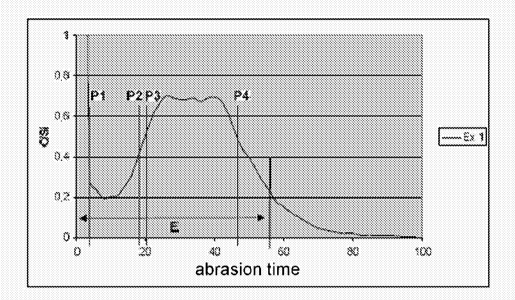


Figure 1

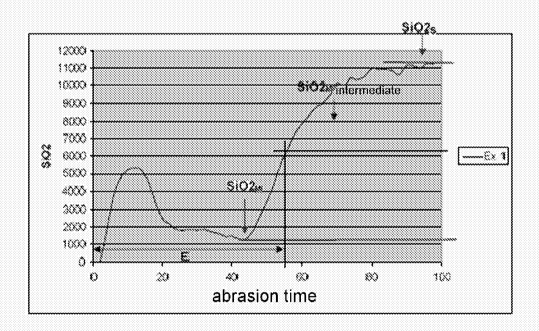


Figure 2

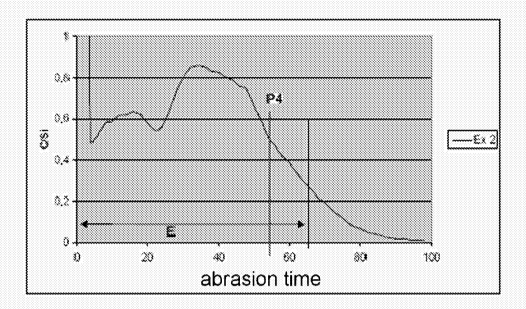


Figure 3