

# An overview of some recent developments in glass science and their relevance to quality control in the glass industry

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**The classical image of glass is that of a rigid, transparent brittle material characterized by a non-crystalline microstructure. This 19<sup>th</sup> and 20<sup>th</sup> century image however is mostly based on the contrast between soda lime glass and metals. It does not really make sense in the 21<sup>st</sup> century where more modern testing methods have increased our understanding of the physiochemistry of glass. Based on recent results and the development of computational molecular dynamic software modelling a new approach to the physiochemistry of glass is outlined. The consequences this view has on glass properties and processing are explained.**

*Keywords: Glass structure, hot working glass, glass processing, effect of glass composition*

## 1 Introduction

Most classical text books on physics, for example the standard A-level physics textbook by Muncaster (1981), divide matter in three types, solid, liquid and gas. Some more recent authors such as Morozov (2012) consider plasma as a fourth type of matter. Glasses however are neither fish nor fowl in this type of division. Many classic physics text, which consider solids from the point of view of crystalline materials, have a problem with glasses. Some of them consider glasses as a super cooled liquid rather than a solid. A good introduction to this long running discussion is given by Curtin (2007).

From an engineering point of view it is strange to consider lead, which flows under pressure at room temperature as a solid, and float glass, which does not flow under pressure at room temperature as a liquid.

From a modern physicists point of view; which accepts that solids can change into other solids or into liquids or gasses, examples being the ferrite-austenitic phase change in iron,

the melting of copper and the sublimation of ice; the rigid categorisation into clear states of matter is dangerous, yet it is still common in both secondary schools and in bachelor programmes of both engineering and natural science. If we take a modern materials science text, such as Callister (2010), chapter 3 deals extensively with the crystalline structure, the next 10 chapters (400 pages) build on this, then in chapter 14 there are 4 pages which deal with glasses before going onto ceramics. In chapter 15 there is some mention of amorphous polymers. So less than 1% of the book is spent on non-crystalline materials, which creates the unplanned impression in students that non-crystalline materials are an unimportant exception.

In general unless one follows some specialised master course in glasses, where typically the book by Shelby (2005) is used, which is the de-facto world standard work on glass; 99+% of engineers and scientists, including those that deal with the processing of float glass on a daily basis typically have no formal training in glass science.

The danger of this is that glass is always viewed from the perspective of metals, specifically a strange solid that becomes liquid at a certain point like steel; and like steel needs to be annealed and can be heat treated. Even the terminology, tempered glass is based on the practice of metallurgical technique. The mind-set that our conventional mind-set favouring metals and other crystalline materials creates, induces an automatic prejudice against glasses and does not provide a scientific or intellectual basis to understand the behaviour of glasses.

## **2 Low and high temperature structure of glass**

The structure of glass is something that cannot be easily determined. X-Ray Diffraction (XRD) only reveals patterns if the material tested is crystalline, which glasses by definition are not. The nano-level structures in glass cannot be seen with conventional microscopes or electron microscopes. Even meso-structures in transparent glasses at the 10 to 100  $\mu\text{m}$  level, such as the bubbles and inclusions in float glass described by Molnar et al. (2013), cannot be seen easily using optical microscopy because there is almost no contrast.

Zachariasen (1932), a noted early 20<sup>th</sup> century crystallographer, published the first scientifically significant model for the structure of glass in 1932. This model is still used even though aspects of Zachariasen's empirical rules for glass formation have been

repeatedly proven wrong in recent decades, as for instance shown by Rao (2002). Huang et al. (2012) proved that the basis concept of the Zachariassen model is true, at least for a 2 dimensional glass lattice. Basically this means that glasses are randomly formed 3d networks of covalently bonded atoms at room temperature. Essentially a single piece of glass is a single macro-molecule. The network can be locally broken up by ionic species, the so called fluxes, which reduce the melting point and improve workability. This however introduces some free local ions in a covalently bonded network with some ionic end incorporated into the covalent network.

Even if this is true at room temperature, this model breaks down when we heat the glass. It is well known that glass can be bend at a certain temperature, for soda lime glass about 800°C, while it melts at a much higher temperature. The problem is that we know nothing about the structure of glass at these temperatures. We do know that as glass is not a metal, the hot bending deformation is not a dislocation based plastic mechanism as we find in metals and which is used in rolling, extruding or bending metals. It is therefore some type of molecular “liquid” flow phenomenon.

As a single macro molecule cannot deform non-elastically, the logical implication is that the glass that we are bending is not a single macro molecule. It must by logic be a series of large molecules bonded together by van der Waals forces. Conceivably the molecules might break down and reconnect during the bending, but nothing is known of the structure or the change of structure of glass at these temperatures, as there are no experimental techniques that allow us to study these things at high temperature.

If we melt the glass again our preconceptions from our studies of metals lead to a dangerous misconception. The macro molecule cannot break down into its constituent atoms. The oxygen molecules would combine and form a gas. The glass would largely sublime rather than melt. The fact that glass exists as a liquid implies that the glass is still composed of molecules in the liquid state. The minor problem is that there is not the single liquid state which we find for metals or in covalently bonded fluids such as water. If we cast glass normally at 1400°C or more the glass is liquid in the sense that it is free flowing, as shown in figure 1, note however the scissors which are necessary to cut the viscous glass stream. If we kiln cast at 950°C the glass is very viscous, flowing like treacle, as in figure 2. Melting and casting takes several hours. At 800°C the glass can be hot bend. In all these three processes the glass is by definition not a single macro-molecule but a molecular

liquid. The difference that determines the viscosity is the molecular weight. There might be dynamic break downs and reconstitutions of the molecules, but this cannot be determined using current experimental technologies.



Figure 1: Casting glass at Poesia



Figure 2: Kiln casting at TU Delft

### 3 Molecular modelling of glass

As there are no experimental tools to investigate the high temperature structure of glass, computational methods have evolved in recent years that allow for the analytical and numerical modelling of high temperature glass structures. Analytically the physiochemistry of glass has been developed into a new model called the topological constraint theory. A good introduction is given by Mauro (2011). Essentially the topological constraint theory explains many reasons why glasses are formed and predicts the glass transition temperature. Figure 3, taken from Mauro (2011), shows the principle of how a glass temperature,  $T_g$ , is calculated. Figure 4, taken from Mauro (2011), shows a ternary diagram and the  $T_g$  prediction related to composition which follows from the topological constraint theory. The predictions from the model have been proven to be quite accurate. Following on from the topological constraint theory is the new and growing field of molecular dynamics modelling of the formation of structures and changes in structure of glasses. This is a numerical technique where atoms are modelled assembling and disassembling into nano-level structures. A good example and explanation can be found in the PhD thesis of Konstantinou (2017).

Konstantinou's modelling allows the prediction of local nano-level structures in glasses. A specific example is how Molybdenum atoms are incorporated into a glass structure

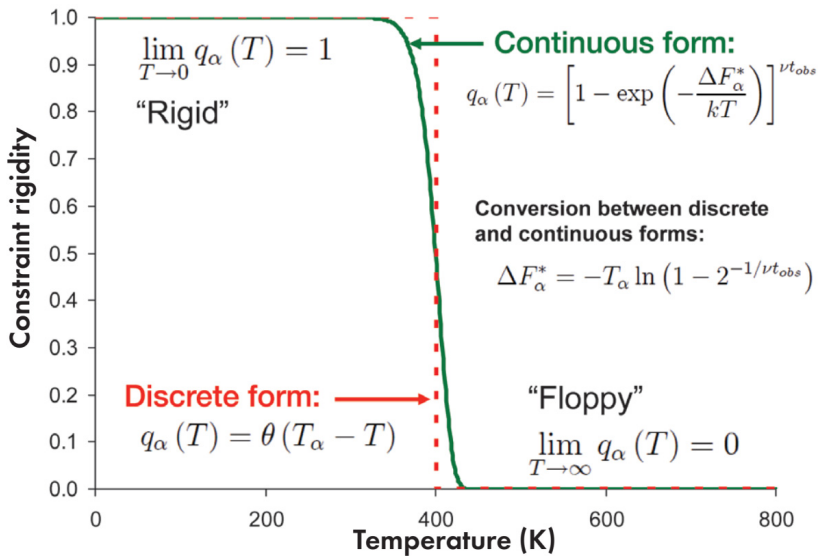


Figure 3: Glass transition predicted from topological constraint model, from Mauro (2011)

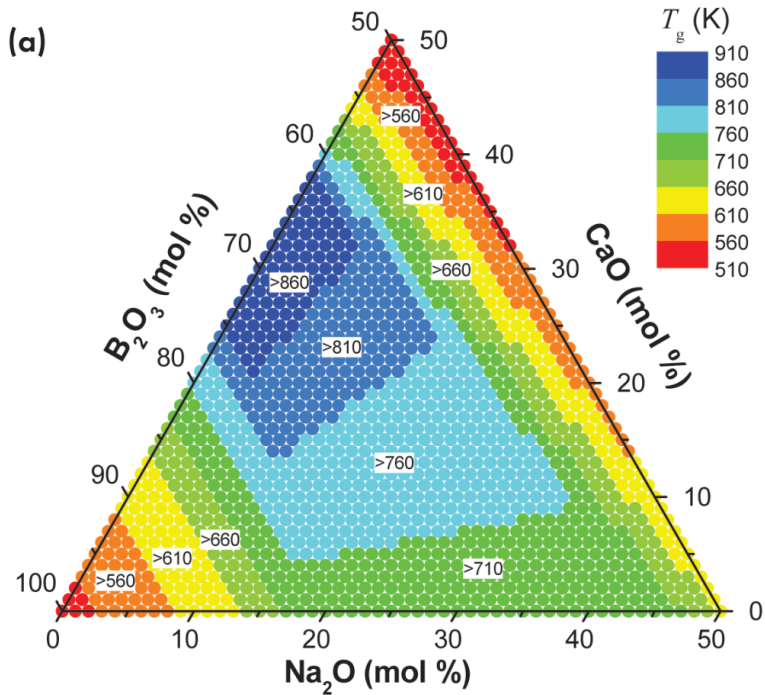


Figure 4:  $T_g$  as a function of composition, from Mauro (2011)

designed for the storage of nuclear materials and is shown in figure 5, with the specific local structure around the Mo atom in figure 6. These modelling techniques allow for close study of the solid glass. Currently exact modelling what happens above the glass transition temperature is beyond the technique but this should be possible within the next 10 years as the technique evolves and more computational power becomes available due to Moore's law. As the strain in the chemical bonds and the strength of the chemical bonds can be calculated, the weakest links in the structure can be identified. It is thus possible to calculate which links break in succession as the temperature increases. Thus the breakup of the macro molecule can be predicted and the resulting development of the molecular liquid modelled. To model the interaction between the molecules is however beyond the current state of the art. Additionally to model the transition of a large enough macro-molecule to several molecules which break down at successively higher temperatures and interact however require a large starting model of the order of 10.000 atoms and preferably more. This is beyond the current capacity of ab initio molecular dynamic modelling.

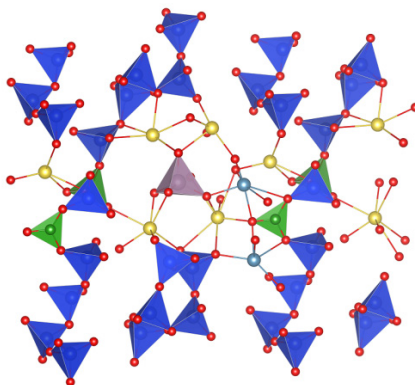


Figure 5: Computed local structure of a silicate glass containing Mo, from Konstantinou (2017)

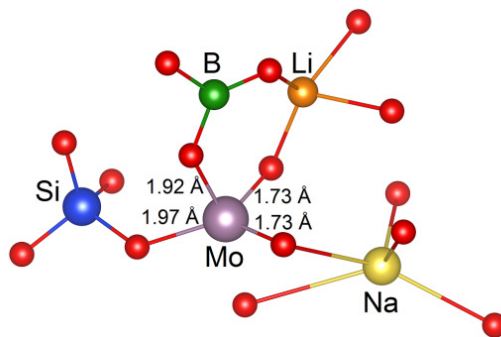


Figure 6: detail of the structure, from Konstantinou (2017)

An important result from both the topological constraint theory and the molecular dynamic modelling is that the glass temperature is shown to be very sensitive to compositional changes, as can be seen in figure 4. The practical relevance of which will be shown later.

#### 4 DSC analysis

The one experimental tool that does work on glasses is Differential Scanning Calorimetry (DSC). In this technique a small sample is heated up and the amount of energy needed to increase the temperature by 1 K is measured. Phase changes and other changes that involve a change in enthalpy are detected in this way, because the increased energy required at the change temperature is measured. In glasses these are commonly used to determine the glass transition temperature,  $T_g$ . Lopes et al. (2014) give a good introduction and example how the DSC technique can be used to study the structure of glasses. DSC analysis has been used to study several float glass samples, such as those in table 3. A result is shown in figure 7. If we compare these to DSC curves in the literature such as those shown in figure 8 which are borrowed from Lopes et al. (2014), some significant differences are observed. The float glass sample shows a minimum enthalpy change for the glass temperature transition. All the other transitions are also less distinct or almost invisible. The float glass sample there has a progressive molecular breakup and no clear transitions. If we do a DSC test on PPG Starphire glass, which is a more expensive quality low iron float glass, using the same settings as before the results shown in figure 9 are obtained. This curve is distinctly different from the curve in figure 7 and a clear  $T_g$  is observed.

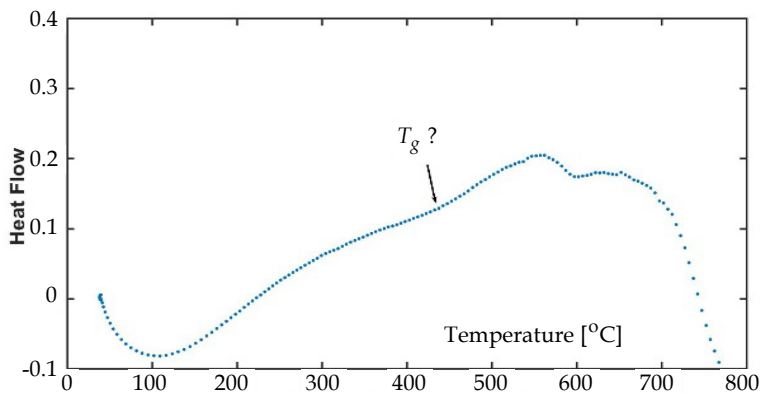


Figure 7: DSC of sample a of table 3

Another important observation can be made from figure 8. Tests with slower heating rates show that the glass temperature seems to shift, similar to that in figure 8. Konstantinou (2017) also concludes the same as the result of his ab initio modelling.

## 5 Actual chemical compositions of commercial glasses

Although there are some norm compositions for normal, mid iron and low iron float glass, these are rarely checked by the end user. In recent years during consultancy work it was necessary to check on the chemical composition of several glass types to explain actual industrial problems. This was done using X-Ray Fluorescence (XRF) analysis. Table 1

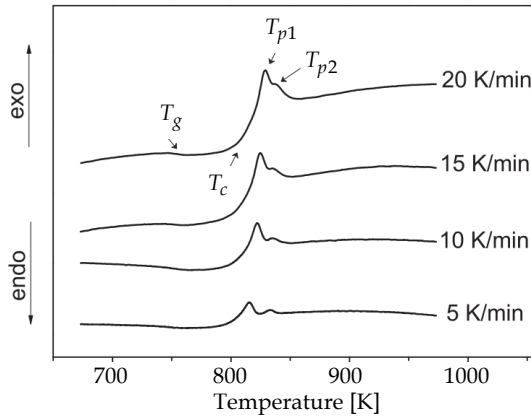


Figure 8: Reference DSC curves from Lopes (2014)

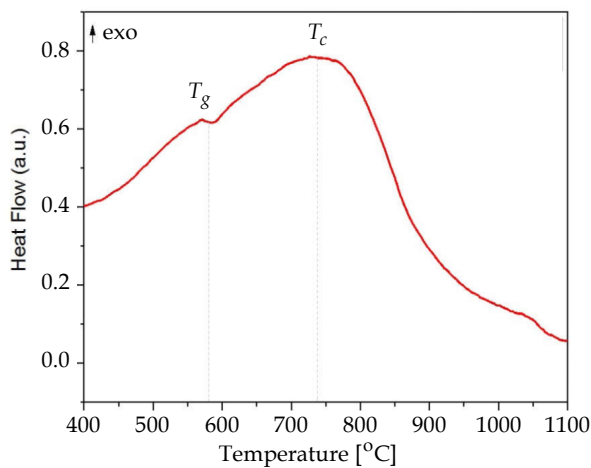


Figure 9: DSC analysis of PPG Starphire float glass, from Bristogianni. et al. (2018)



contains the results of 3 samples of 1 mm thin glass supplied to a manufacturer of a specific product, involving 3 different factories and 2 producers. Table 2 contains the results of one type of bulk coloured glass with multiple samples produced at a single factory at different times. Table 3 contains the results of several types of float glass which a single processor bought from different European suppliers and also contains the norm composition for float glass as a reference. It is clear that the compositions differ significantly. Table 1 shows that different factories even from the same owner have different compositions. Limiting the glass that was bought to a single factory, in this case producer A factory 2, significantly reduced the failure rate during production of the product. This reduced the drop-out rate significantly making the product much more economical to produce. Table 2 shows that the chemical composition of a single float line is reasonably constant, although there is a clear variation in time. Table 3 shows that many modern float glasses have compositions that are modified for lower melting temperature to reduce the cost. This will also affect the temperatures for correct processing of the glass for tempering and bending. There are also considerable differences between different suppliers. As glass which is bought by a processor is not tested in terms of composition, there is no guarantee that the glass is processed at the optimum temperatures. All glass that is used is assumed by the end user to be the same and is processed the same, irrespective of whether a supplied batch needs slightly different settings. In practice most end users do not know where the glass they buy was produced and if they buy from different sources the glass is routinely mixed up in the store house. The methodological way of checking and inventorying materials in the aerospace industry is the complete reverse of how the glass industry treats its materials with resulting differences in quality control and consistency.

*Table 1: Chemical composition of 3 types of thin glass in wt. %*

Compound	Producer A factory 1	Producer A factory 2	Producer B
SiO <sub>2</sub>	74.83	73.84	74.36
Na <sub>2</sub> O	12.84	12.83	12.25
CaO	7.24	7.17	8.30
MgO	4.29	4.35	4.07
Al <sub>2</sub> O <sub>3</sub>	0.82	1.24	0.48
Fe <sub>2</sub> O <sub>3</sub>	0.023	0.077	0.019
ZrO <sub>2</sub>	0.005	0.008	0.006

Table 2: Chemical composition of bulk glass produced at different times in one factory, wt%

Compound	Float glass composition by norm	Sample a	Sample b	Sample c
SiO <sub>2</sub>	71.9	74	72.7	74
Na <sub>2</sub> O	13.1	12.8	12.3	12.6
CaO	9.23	7.7	7.7	7.9
MgO	5.64	4.1	3.9	4.1
Al <sub>2</sub> O <sub>3</sub>	0.008	0.7	0.5	0.4
Fe <sub>2</sub> O <sub>3</sub>	0.04	0.52	0.50	0.62
K <sub>2</sub> O	0.02	0.06	0.12	0.11

Table 3: Chemical composition float glasses produced by different factories, wt%

Compound	Float glass composition by norm	Sample a	Sample b	Sample c	Sample d
SiO <sub>2</sub>	71.9	64.72	68.54	66.59	70.41
Na <sub>2</sub> O	13.1	12.61	12.35	12.41	12.89
CaO	9.23	16.83	13.20	15.67	11.31
MgO	5.64	4.00	3.82	3.94	4.43
Al <sub>2</sub> O <sub>3</sub>	0.008	0.79	0.68	0.71	0.02
Fe <sub>2</sub> O <sub>3</sub>	0.02	0.18	0.15	0.22	0.19

## 6 Effect of composition on glass viscosity

In glasses the various working temperatures are related to the viscosity. Martlew (2005) gives these as:

- Melting point 2.0
- Working point 4.0
- Flow point 5.0
- Littleton's softening point 7.65
- Deformation point 11.5
- Annealing point 13.0
- Strain point 14.5

The values are the 10 base logarithm of the viscosity in poise. 10<sup>2</sup> means 100 poise, etc.

These are however completely arbitrary definitions and based on the practices of

traditional glass working. Actual values for the different viscosities are difficult to find. Martlew (2005) gives an incomplete series of values for different groups of glasses. For Na<sub>2</sub>O -SiO<sub>2</sub> melts the data is summarized in table 4. Increased Na<sub>2</sub>O content decreases the melting point, from 10 to 15% by 32 K per mol% increase. From 25 to 30% by 15.6 K per mol% increase. The other glass working temperatures are similarly affected.

*Table 4: Melting point as function of composition for Na<sub>2</sub>O SiO<sub>2</sub> melts*

Na <sub>2</sub> O mol%	SiO <sub>2</sub> mol%	Temperature °C
10	90	1728
15	85	1568
20	80	1455
25	75	1372
30	70	1294
35	65	1205
40	60	1015
45	55	1057

## 7 Discussion

In the artistic glass world there is a significantly greater appreciation of the different behaviour of different glasses. This arises simply from the need to work many different glasses into complex objects. Their publications are however not widely read in the general glass community. Stone (2000) gives for many types of glasses a range of temperatures. Stone (2000) also indicates a clear need to be cautious in using these values. To quote several relevant passages from p 6.28 en 6.29 from Stone (2000):

“The most significant variable after thickness is the glass type being fired. Glasses differ in their heat cycle requirements in four important areas:

1. the rates of temperature change, both up and down
2. the amount of annealing time
3. the annealing temperature
4. the subsequent down phase temperature settings.”

“The following specific temperatures for the annealing stage are approximate, especially regarding glass from producers that supply a range of colours and forms.”

“Remember that glass has an annealing range rather than a single point where stress is relieved. It’s just as well, because not all manufacturers test the temperature at which the viscosity of their glass is  $10^{13}$  Poises, the accepted measurement of an annealing point.”

These points derived from significant practical experience in the artistic glass community, do however translate into problems for the current float glass industry. If we look at the most important quality control problems in the float glass industry:

1. uneven tempering
2. optical anisotropy
3. roller wave distortion

The first of these have been studied by Chen et al. (2013) and Veer et al. (2016). Tempered float glass is actually quite inhomogeneous in terms of the surface pre-stress. The surface pre-stress varies widely in a single specimen and is also highly variable between specimens. Although Chen et al. (2013) correctly attribute this result to uneven cooling, different physiochemical response within single float glass specimens and between specimens with different compositions cannot be ruled out. As the glass point varies with changes in compositions and is also dependent on the heating rate and cooling rate; equal treatment of chemically different float glass panels will result in different compressive pre-stresses. Additionally inhomogeneous cooling rates will affect the glass temperature locally of the glass. The different compressive pre-stresses cause visual anisotropy because the different compressive pre-stresses cause different local polarisation effects which cause blotchy patterns. This effect and the problems it creates are described quite well by Pasetto (2014) and are shown to be a significant problem in the current glass industry. The lack of internationally or even nationally accepted quality control norms regarding optical anisotropy of course does not help. There is some “reliable” anecdotal evidence that some Chinese glass producers that single source their float glass from a close-by factory and optimise the settings of their tempering furnaces for this glass have (significantly) less problems with an-isotropy. Although this would argue for a role of composition effects, anecdotal evidence is not however scientific evidence. It is however clear from discussions

with façade sub-contractors that some producers have a much bigger problem with anisotropy than others.

Roller wave distortion is another critical quality control problem, again beset by a total lack of standards in the industry for what is acceptable or even a standardised way of measuring it. A good overview of this particular problem is given by Abbot and Maddocks (2001). There is however a clear relation between roller wave distortion and a processing temperature which is too high for the actual glass being tempered. Abbot (2001) states specifically that for tempering:

“In theory, heat treatment requires uniform heating of the glass to  $621 \pm 3$  deg C, while holding the glass in a flat state”

This might be true for a certain float glass, for instance the PPG Starphire from figure 9, it is very doubtful if this is correct for the glass from a different supplier in figure 7, where the  $T_g$  is clearly lower and is also not very well defined.

As it is clear from the previous paragraphs that the glass temperature is dependent on the heating rate and that glass producers who do not single source their glass, by necessity will have to deal with different chemical compositions and thus (slightly) different glass temperature. Some glass panels will inevitably be tempered at too high a temperature. A temperature which is too low is less likely as this requires a higher melting glass which is more expensive to process, which is what the industry does not want. As figure 4 and table 4 show, a couple of % difference in one or two components of the glass can shift the glass transition by tens of degrees centigrade. The resultant viscosity changes can contribute significantly to roller wave distortion. The differences in composition in table 3 are large enough to cause the differences in  $T_g$  that would significantly increase the risk of roller wave distortion. Some of these glasses also have an indistinct  $T_g$  point as shown by DSC measurements, such as in figure 7, which makes it difficult to determine the correct processing temperature.

Quality control in the glass industry is thus more than proper cutting and grinding, and setting the right temperature settings on the tempering furnaces. It needs to either single source the glass from a single factory and optimize the machinery for this glass or adapt the settings continuously to the glass being processed at that moment. Table 2 shows that

glass from a single float line is reasonably consistent in terms of composition which should allow for a single set of processing temperatures. Or the industry needs to adjust the process settings to the actual chemical composition of the glass being processed.

However the current logistics of the industry do not allow for differences in composition other than normal float, mid iron and low iron glass. This is also because the effect of relatively small differences in composition are not considered as important. If we compare this to the metallurgical industry where exact compositions are controlled and the logistics setup allows for streams of materials being correctly controlled allowing for differences in composition and in different heat treatments there is much to learn for the glass industry.

## 8 Conclusions

From the previous paragraphs the following conclusions are drawn:

- There are significant differences in the float glass composition if we look at glass from different suppliers.
- These differences are sufficient to shift the glass temperature potentially by 10 K or more as shown by both DSC tests and modern analytical and numerical modelling.
- These differences can contribute to industrial problems such as uneven tempering, optical anisotropy and roller wave distortion.
- Quality control in the glass industry should allow for compositional differences.

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