

Experimental mixing of natron and plant ash style glass: implications for ancient glass recycling

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ABSTRACT

The practice of re-melting glass was well known, certainly from the Roman period onwards. This can be seen not only in ancient literary evidence but also in the archaeological evidence, collections of broken glass have been found in, for example, Pompeii (79AD) and the Iulia Felix shipwreck (3rd century AD). Elevated levels of certain transition metals in archaeological glasses are interpreted as indications of the mixing and/or recycling of different glasses. Assumptions have been made that all glasses could be recycled, but to what extent are these valid? Why does the evidence for the recycling of glass only occur from the Roman period onwards? From the middle of the 1st millennium BC to the 9th century AD, natron glass was the predominant glass type in the Mediterranean and Europe, however, plant ash glass was still in use in some areas. To test the effects on the final product of mixing different composition glass types, experimental glasses were made by mixing varying quantities of replica plant ash glass, replica natron glass, and a modern glass. At low temperatures crystalline material formed in the products containing replica plant ash glass. As the plant ash glass content increased, so too did the amount of crystalline material produced. This is due to a combination of the glass compositions and the firing temperature. It appears that natron type glass can be more easily recycled at lower temperatures, although, if a high enough temperature is used then most glass types can be recycled. Early furnace technology, i.e. the vertical heating chamber furnace, may not have been able to achieve these high temperatures, hence the widespread practice of recycling did not begin until after the invention of glassblowing which required a change in the furnace technology to the use of a horizontal heating chamber furnace.

KEYWORDS

Recycling; Plant Ash glass; Natron glass; Devitrification; Crystallisation

INTRODUCTION

The concept of recycling glass has long been taken for granted¹, although the exact date of its discovery is unknown². The literary evidence for recycling suggests that the discovery took place sometime in the Flavian period (1st century AD)². Writers such as Martial (*Epigrams* 1:41; 10:3) and Juvenal (*Satires* 5) mention the trade of broken glass; while Pliny the Elder (*NH* 36.67) says that glass can be stuck back together but never completely fused again. However, the idea that glass could be completely re-melted, did reverberate widely in Roman literature^{2,3}. Recycling glass would have involved (re)melting waste products from glass workshops and/or old or broken glass¹. The assumption that fragments of glass or cullet could be easily added to the batch has provided the basis for a number of theories seeking to explain the composition of Roman glass^{1,4-13}. It has also been suggested that the manufacture

of primary glass (i.e. made from raw materials) required more skill and knowledge than the production of secondary glass (i.e. worked glass), since the glass worker need only re-melt raw glass and/or cullet¹⁴. Conversely, the technical knowledge required to ensure the correct viscosity of the glass is probably one of the most important skills in glass shaping^{15,16}.

Collections of Roman and post-Roman broken glass or cullet have been found on a variety of archaeological sites, including shipwrecks such as the 3rd century AD Iulia Felix^{11,17} and in secondary glass workshops including at Bet She'an, Israel (6th-7th century) and in the 1st century BC Jewish Quarter of the old city of Jerusalem¹⁸. A basket of scrap glass was found at Pompeii, which confirms that the collection of broken glass occurred prior to 79AD^{1,19}. Finally, the chemical compositions of archaeological glasses have been used to suggest that recycling, or the reuse of old glass occurred^{1,20-26}. This is proposed based on elevated levels of certain trace elements (Co, Cu, Mn, Mo, Ni, Pb, Sb, Zn...) within the glass^{4,7,20,21}, and on the use of isotopic mixing lines¹. Elevated levels of trace elements are defined as above the 'natural' level found in the raw materials²⁷, but not high enough to constitute a deliberate addition to the glass melt. For example, if colourless glasses were being recycled and a small amount of coloured material was accidentally incorporated into the batch, this would have the effect of elevating the associated colouring elements without impacting on the colour of the glass.

Roman glass is soda-lime-silica in composition, with the majority being made from natron, a mineral soda. The use of natron glass spans the whole Roman period but was not solely limited to this era, being manufactured between the mid-first millennium BC and the 9th century AD. It is characterised by low magnesium and potassium contents (MgO and K₂O less than c.1.5%) and is sometimes referred to as LMLK glass^{7,28-30}. A 'typical' natron glass can contain 54-74wt% SiO₂, 3-8.5wt% CaO, 13-20wt% Na₂O and around 2.5wt% Al₂O₃³. Prior to the mid-first millennium BC, a different form of flux was used in glass to lower the melting temperature of the silica. In this case, the soda content of glass originates from the use of plant ashes. Their production dates back to at least the late Bronze Age in Egypt and Mesopotamia. Due to the varying nature of the plants used in their production, their chemical composition is variable. An example of Egyptian plant ash composition glass is that from Amarna, dated to the 14th century BC. This glass contains on average 64.27wt% SiO₂, 1.70wt% Al₂O₃, 3.98wt% MgO, 8.30wt% CaO, 18.38wt% Na₂O, 0.63% Fe₂O₃ and 1.44wt% K₂O³¹. After the 9th century AD, there is a switch back from natron to plant ash glass, either soda-rich in the Eastern and Southern Mediterranean area, or potassium-rich in Northern Europe.

The periods of use for specific types of flux and glass are general. There would have been a certain amount of overlap between the uses of plant ash as a flux and natron. Also, not all glass manufacturers may have switched to the use of natron, some may have preferred to continue using plant ashes as a soda source³², for example, the Sassanian glasses produced between the 3rd and 7th centuries AD. The recycling and reuse of glass clearly occurred from the Roman period onwards. However, assumptions have been made that all glass could be recycled. This research aims at testing this assumption. What, if any, effects would the mixing of different glass compositional types have on the final product? Three different compositional types of glass, replica plant ash, replica natron glass and a modern glass (representing a generic mineral soda glass) were mixed and the chemical composition of the resulting glasses were evaluated. The mineralogical composition of devitrified, crystalline phases was also determined.

METHOD

A replica plant ash recipe glass, i.e. a glass that had been made with modern lab grade raw materials representing a glass made with plant ash (16.8wt% Na₂O, 8.0wt% MgO, 1.0wt% Al₂O₃, 62.3wt% SiO₂, 2.4wt% K₂O, 8.5wt% CaO, 0.4wt% Fe₂O₃), and a replica natron recipe glass (17.4wt% Na₂O, 1.7wt% MgO, 2.5wt% Al₂O₃, 70.8wt% SiO₂, 0.8wt% K₂O, 7.0wt% CaO) were powdered and mixed together. These replica glasses only contained the major and minor elements present in the glass, common impurities such as Cl or S were not added. A third, modern, colourless glass, (15wt% Na₂O, 3.2wt% MgO, 1.1wt% Al₂O₃, 73.4wt% SiO₂, 6.3wt% CaO), was also used. The modern glass is similar in composition to a mineral soda glass, with the exception of showing higher amounts of MgO. This is because modern manufacturers add small amounts to protect against devitrification³³. Table 1 shows the varying percentages of each glass that were mixed together to create the mixed and re-molten glasses. Each glass batch was placed in a mullite crucible and then into a muffle furnace, where the temperature was raised to 900°C. The melts were then held at this temperature for 4 hours before being allowed to cool down in the furnace. A temperature of 900°C was chosen because secondary glass working furnaces were operated at lower temperatures compared to primary furnaces, depending on the primary function of the furnace¹⁶. A calculation of the working and softening temperatures (<http://glassproperties.com/>) for each of the glass mixes used in this work indicated that 900°C fell within working range for all of the glasses.

The compositions of the resulting glasses were determined by ICP-OES. Samples were ground to a fine powder using an agate mortar. 100mg was weighed and dissolved by alkaline fusion with 500mg of LiBO₂ at 1000°C in graphite crucibles. The fusion was then poured into 50ml of 0.43M HNO₃ in a polypropylene beaker. The solutions were further diluted (10-fold) by adding 0.43M HNO₃ before analysis. The compositions were determined using a Varian 720-ES equipped with a SeaSpray concentric glass nebuliser, a double-pass cyclonic glass spray chamber and a 'high solids' torch. The instrument features a Cooled Cone Interface, echelle monochromator and custom-designed Vistachip CCD detector mounted on a triple-stage Peltier device and cooled to -35°C. Solutions were presented to the spectrometer using the Varian SPS3 Sample Preparation System. Table 2 lists the operating parameters used for the ICP-OES analysis. The oxides determined were Na₂O, MgO, Al₂O₃, SiO₂, K₂O, CaO, TiO₂, Fe₂O₃, BaO. Calibration curves were created using solutions of four silicate-rock reference materials (GBW-7114, BR, BCS-CRM 267, BCS-CRM 269) and four glass reference materials (NIST SRM 610, NIST SRM 612, NIST SRM 620, SGT 10). Blank solutions were created by mixing the same ingredients and following the same procedure as used for the sample dissolution, but without the glass powder. BR was measured every 7 samples in order to check for possible drift of the instrumental response. Results were accurate to within 10% of the certified values for most major and minor elements, with the exception of MgO, TiO₂ and Fe₂O₃, these latter three were all within 20%.

Some of the resulting glass samples appeared to have crystalline material present, therefore, it was decided to perform qualitative XRD analyses. Powdered samples were micronized in a McCrone Micronising mill for 5 minutes with 5ml ethanol as a grinding agent. After two days of drying, the samples were disaggregated in a mortar and passed through a 250µm sieve. The samples were measured with a Phillips PW180 diffractometer with a Bragg/Brentano θ -2 θ setup and CuK α radiation at 45kV and 30mA. The scan range was from 5-70° with a step size of 0.02 and 4 seconds per step. The software used for peak identification was EVA. Thin sections were also made of each sample.

RESULTS

Table 3 indicates the results of the ICP-OES analyses of the samples. The ‘recycled’ glasses all form mixing lines between the parent compositions, with the exception of Na_2O . In the case of the two glasses made as a result of mixing all three parent glasses, these sit within the triangle formed by the parent glasses. When the samples were removed from the furnace, all had a blue-green colour due to the iron in the original ‘plant ash’ glass. Several of the samples appeared crystalline in nature (Figure 1). XRD analyses indicated that sample RBS-D, made from 75% plant ash glass and 25% natron glass, had three mineral phases present quartz, combeite ($\text{Na}_2\text{Ca}_2\text{Si}_3\text{O}_9$), augite ($((\text{Ca},\text{Na})(\text{Mg},\text{Fe},\text{Al})(\text{Si},\text{Al})_2\text{O}_6)$) with the possible presence of a fourth phase in the form of pseudowollastonite (CaSiO_3). The other two plant ash/natron mixed glasses contained only three definite mineral phases (augite, quartz, combeite). In the samples made from a mix of all three parent glasses, only two mineral phases were observed (augite, quartz). A study of thin sections made from all of the samples indicated that as the amount of plant ash glass increased, so too did the amount of crystalline material in the glass (Figure 2).

DISCUSSION

Figure 2 shows that as the amount of plant ash glass is increased, the amount of crystals in the resulting glass also increases. This is a function of the glass chemistry in combination with the firing temperature used to remelt the glasses. The natron parent glass had 17.4wt% Na_2O , and 7.0wt% CaO , whereas the plant ash parent glass had 16.7wt% Na_2O and 8.9wt% CaO . The resulting mixed glasses show a loss of Na_2O , compared to the predicted values (Table 4). This loss of Na_2O compared to the amount of CaO present is not great enough on its own to cause the devitrification of the glass. However, the glasses also contain MgO , and varying the amount of this component can have dramatic effects on both the liquidus temperature (the temperature above which all material is molten) of the melt and the subsequent formation of crystalline phases³⁴. Likewise, varying the amount of K_2O will change the viscosity of the glass and the subsequent ability of the melt to form crystalline phases³⁵. By using a liquidus temperature calculator (<http://glassproperties.com/>) it was possible to determine the approximate liquidus temperature for each of the glasses used in the experiment (Table 5). The calculator models the temperature based on the common major and minor components found in soda-lime-silica glasses (Na_2O , MgO , Al_2O_3 , SiO_2 , K_2O , CaO) and compares the entered values to glass from the SciGlass (www.sciglass.info) database. The database glasses all had the liquidus temperature determined experimentally. The calculated values are, however, only approximations because the SiO_2 component of the glass is calculated and cannot be entered manually. These estimations show that as the amount of plant ash glass incorporated in the mix increases, so too does the liquidus temperature of the melt. It should be remembered, however, that the liquidus temperature is the point at which the total mix is molten. At lower temperatures different proportions of the batch will be in a liquid form, depending on the composition of the glass^{36–38}.

A decrease in the temperature needed to melt the glass implies a weakening of the silicate network and a decrease in the viscosity of the glass³⁹. In terms of glass working and the manufacture of objects, this is arguably a good thing, since it would require less fuel to operate the furnace. However, this weakening of the network increases the risk of crystallisation of the melt³⁹. In order for crystals to form in the melt, crystal nuclei need to be present⁴⁰, and these need to grow at a suitable rate⁴¹. Since the samples in this experiment were formed from existing glasses it is unlikely, although not impossible, that residual crystal

nuclei were present in the originals. What is more likely is that the powdered nature of the samples prior to heating acted as nuclei for further crystal growth⁴². Since powdered particles would have been present throughout the melt and the liquidus temperature was never reached, this would explain the presence of crystal growth throughout the bulk of the samples. These results clearly show that the quantity of the crystals forming in the glasses is proportional to the amount of plant ash glass present. This is because as the amount of plant ash glass increases, the distance between the liquidus temperature of the glass and furnace temperature used increases. A study by Bingham and Marshall⁴³ suggested minor alterations to the composition of a glass can have large effects on changes to the viscosity. They defined a key parameter in the determination of crystal formation as the difference between the 'forming' and liquidus temperatures (ΔT_{FL}) in the glass. The 'forming' temperature is the temperature at which the glass begins to be shaped into an object. As ΔT_{FL} decreases into negativity, so the risk of crystallisation increases. The forming temperatures, or the temperature at $\log 10^4$ Pa.s (i.e. the temperature of the melt at a glass forming viscosity), for the experimental glasses were calculated using a viscosity calculator (<http://glassproperties.com>). The subsequent ΔT_{FL} for each sample was calculated (Table 5); this method also clearly shows that while the natron-rich glasses are not immune to crystallisation, the plant-ash-rich glass mixes are more likely to devitrify. XRD analyses of glasses mixed from only replica natron and modern glass indicated that no crystalline phases were present in these mixes, despite the negative ΔT_{FL} . Some modern manufacturers give anecdotal reports of operating with negative ΔT_{FL} without any adverse effects, but this may be due to their furnace conditions and forming operations⁴³.

It should be noted that glass working occurs at a range of temperatures and viscosities³⁸ because, for example, glass blowing happens outside the furnace. The initial 'working' viscosity ($\log 10^3$ Pa.s) is the point at which the melt is the correct viscosity to be gathered as a gob ready for glass forming. However, the viscosity of the glass varies during formation processes¹⁶. For example, the viscosity at the surface of a bottle during blowing was measured as being between $\log 10^5$ Pa.s and $\log 10^9$ Pa.s³⁸. The working range of the glass is then described as being the working temperature minus the softening temperature. The smaller this number is, the shorter the working range of the glass. The temperatures needed to achieve these various viscosities are in turn dependent on the composition of the glass. While a knowledge of the working range of a glass is useful for estimating the time available for forming processes, it does not inform about the crystallisation resistance of the glass³⁸. This is determined by the time and/or cooling rate of the melt below the liquidus temperature³⁸. The ΔT_{FL} is a good indicator for the devitrification potential of a glass. However, for the study of ancient glass, the viscosity of the specific formation method, i.e. blowing, sagging, casting etc, should be used in place of the generic forming temperature.

Crystals form in a glass melt during cooling and any vitreous material which is heated for long enough above its glass transition temperature (T_g) can crystallise⁴⁴. The T_g is the vitrification point of the melt; below the T_g the material is a glass, above the T_g it is molten. Nucleation of crystals usually occurs just above T_g , after processing and during cooling to a glassy state⁴¹. The nature of the crystals formed depends on the composition of the melt. The two samples which were mixed from three parent glasses had two crystalline phases (augite, quartz), the 75% plant ash and 25% natron glass had three to four crystalline phases (augite, quartz, combeite, pseudowollastonite). The remaining glass mixes had three phases (augite, quartz, combeite). Generally small additions of MgO and Al₂O₃ in a glass will help prevent devitrification^{33,42}. Too much MgO or Al₂O₃ will, however, promote crystallisation⁴⁵. Where crystals form, there is a resulting change in chemistry of the melt, depending on which

components become bound to the structure of the crystals⁴². For example, silica-rich and/or alkali or alkaline-earth rich regions become more apparent⁴⁶. This can lead to new crystal phases being formed through the redistribution of cations in certain areas of the silicate framework, which adjusts to their presence⁴².

Most of the work looking at crystal formation in archaeological glasses has occurred from the perspective of creating glass from raw batch materials rather than re-melting existing glasses. In these cases, the crystals were formed during batches fired for short periods of time and were forming from the re-crystallisation of undissolved particles in the melt⁴⁷. The limit of solubility of a given element within a glass melt is governed by the temperature, therefore, crystallisation of suitable phases occurs when these limits are crossed⁴⁸. For example, the maximum amount of CaO which can be incorporated into a typical Late Bronze Age glass was found to be c. 6wt% at 900°C and c. 10wt% at 1000°C⁴⁸. Previous studies have also shown that the maximum amount of MgO that can be incorporated into a glass melt before diopside ($\text{CaMgSi}_2\text{O}_6$) crystallisation occurs is c.4-5wt%^{48,49}. Sample RBS-D which contained 75% plant ash glass and 25% natron glass and RBS-PA which was a re-melt of 100% replica plant ash glass were the only samples to potentially form pseudowollastonite during cooling. However, all the glasses mixed with plant ash glass formed quartz and augite. RBS-D has 7.96wt% CaO and 6.08wt% MgO. Therefore, at 900°C, there would have been an excess of both CaO and MgO in the melt. Since augite formed, rather than diopside, this means that Na, Al and/or Fe were also incorporated into the structure.

Studies of commercial soda-lime glasses (15.90% Na_2O ; 0.52% K_2O ; 3.63% MgO ; 5.13% CaO ; 2.17% $\text{Al}_2\text{O}_3 + \text{Fe}_2\text{O}_3$; 72.23% SiO_2) found quartz forming in the melt at around 862°C³⁷. Similarly, Volzone and Stábile⁵⁰ found a form of combeite occurring at c.800-900°C. Combeite predominantly contains Na, Ca and Si, but it too can incorporate Al and Fe into its structure. The pseudowollastonite is harder to explain, since this is usually formed at higher temperatures⁵¹⁻⁵⁷. However, Mitsuda and Banno⁵⁸ accidentally formed pseudowollastonite from a mix of lime and quartz at around 200°C. The combination of quartz crystals and excess CaO availability may have led to the potential production of pseudowollastonite. However, as the amount of plant ash glass is reduced, the viscosity of the resultant melt is increased, thereby making the formation of pseudowollastonite harder. The increased viscosity of the melt means it is harder for the cations to redistribute and form new crystalline phases. The CaO is then incorporated in other mineral phases rather than in pseudowollastonite.

It is important to note, however, that it can be extremely difficult to predict which devitrification phases will be present in a given glass, since this is dependent on the composition of the original glass, the nature of the furnace and crucibles used, the chosen furnace operating temperatures, and the speed with which the melt is subsequently cooled^{37,51,59,60}. Trace level impurities in the glass can also affect the viscosity of the melt and subsequently impact on the rate of devitrification⁶⁰. Likewise, increasing a minor oxide in the glass may result in the precipitation of a multi-element phase⁵⁹. In essence, if a glass melt is held at a temperature favourable to crystal development for long periods of time, then this glass is likely to devitrify. This temperature changes depending on the composition of the melt³⁵.

Archaeological Implications

Previous work, which compared natron and plant ash glass compositions, found that the earlier plant ash glasses formed around the soda-rich cotectic. The silica-rich cotectic is favoured by natron glass⁶¹. This means that plant ash glasses can be made using shorter firing times with less fuel requirements⁶¹. In terms of the current research, where the focus is on the mixing of different glass compositions, the results clearly show that natron composition glass can be remelted more successfully than plant ash glass at 900°C. From the work of Shugar and Rehren⁶¹, Van Beeumen⁶², Bingham and Jackson⁶³ and others, it is clear that Roman primary glass furnaces would have had temperatures capable of re-melting all types of glass cullet. However, the success of this would be partially dependent on the amount of cullet used and what proportions of that cullet were of different compositions. As Freestone²⁸ suggests, if glass recycling occurred on an intra-regional basis, then the glasses being mixed were likely to be of the same basic type. The incorporation of plant ash glass cullet to a natron type raw batch would almost certainly result in a degree of crystallisation. However, if only a small amount of plant ash cullet were used, then it may have been possible, if necessary, to pick out the devitrified pieces when the raw glass was broken up. The higher temperatures in the majority of the tank furnace may also have resulted in less devitrification, but this remains to be modelled. It is more likely that crystallisation would still occur, but the nature of the crystals may vary, for example, at between 800°C and 1100°C wollastonite (βCaSiO_3) is known to form⁶⁴. The absence of wollastonite in our samples is due to the low temperature used for firing, although pseudowollastonite phases did occur as the plant ash glass content increased.

In terms of secondary glass working, natron glass could be re-melted and recycled either on its own or as part of a mixed batch. The nature of the working furnace would have played a significant role in terms of whether the glass could or could not be successfully recycled. For example, the early furnaces with a vertical heating chamber achieved a lower temperature, whereas the later horizontal heating chamber furnaces allowed glass workers to use molten glass². At lower temperatures, mixing glass compositions may have created some problems in terms of devitrification and the removal of bubbles, but it would certainly have been possible to recycle the glass. The main crystalline phases forming in the three glass mixes were augite and quartz, but of this, the amount of crystalline material was greatly reduced when the amount of incorporated plant ash glass was reduced. Freestone et al⁶⁵ analysed some Anglo-Saxon period glasses which appeared to be a mix of both natron and plant ash material. They concluded that these 6th century AD glasses were more likely natron glasses with the addition of a quantity of plant ash material to extend a restricted supply of glass, rather than the mixing of natron glass and plant ash glass. However, around the 6th century AD, attempts could have been made to extend the available natron glass by the addition of up to 10% plant ash glass⁶⁵. It should be noted however, that the composition of the later plant ash glasses would be different again to that used in this study. Therefore, even if devitrification did occur when natron glass was recycled, it probably happened to a relatively small degree. The glass mix itself could still be worked although the final glass product may have had flaws. *“Like the Roman glassblower’s furnace, furnaces in Herat (Afghanistan), Damascus, Hebron, and Cairo still functioning in the 1960s and 1970s remelted broken glass, but the quality of their output was poor. The glass is bubbly and full of striae and other impurities, in part because they did not achieve the high temperatures required for complete fusion”* (Stern, 1999: 451-452).

Late Bronze Age plant ash glass would have been difficult to recycle in secondary furnaces. In very small quantities, this may have been possible, but would probably have been detrimental to the final product. It could have been possible to refine the results of plant ash-

rich glass mixes and improve the amount of glassy material collected (as has been demonstrated in other research ⁶¹), but this would have required time, fuel and resources. What may be more probable is that small amounts of plant ash glass were, if at all, recycled as cullet added in the glass making stage, where fresh mineral matter and plant ashes are molten. The composition of the early plant ash glass is such that it could not be worked and re-melted in the same way as natron glass. The increased K_2O in plant ash glass will increase the viscosity of the melt and thereby reduce the workability of the glass ³⁵. An important modern observation on the use of cullet in glass manufacture is that as the composition of the cullet varies, so too will its effect on the final glass. Therefore, batch calculations are necessary to compensate for the compositional difference ⁴³. Late Bronze Age and Early Iron Age glass was also strongly coloured. Therefore, unless the glass was well sorted, any attempt to remelt it would probably result in a dark or black glass. Early attempts to completely re-melt glass may have failed and/or resulted in a devitrified product, possibly even with an unappealing colour. This in turn would have led people to believe that glass cannot be re-melted or reused. As natron glass came into use, waste from glass manufacture could be included and reused, encouraging people to again try recycling broken glass.

Late Bronze Age plant ash glass was prone to devitrification when re-melted. The increased risk of crystallisation coupled with the shorter working ranges of plant ash glass could indicate why the invention of glass blowing and recycling may not have occurred until after natron glass became the predominant type. The subsequent use of horizontal heating chamber furnaces, coupled with the variation in plant ash glass compositions, meant it eventually became possible for plant ash glasses themselves to be recycled. This is evidenced by the analysis of glasses from later periods, where sodic and potash glass has been mixed and recycled ^{32,66}. The discovery that glass could be recycled would, as Stern ² puts it, have been revolutionary. Yet, the speed and extent to which this new idea spread would have been dependent on political, economic and local factors. Since a small change in the composition of the glass could potentially have had a very dramatic effect on the properties and workability of the glass, artisans may have been cautious. In areas where only one type or composition of glass was in use, the concept of recycling may have spread very quickly. The benefits of recycling old glass, such as the reduction of fuel consumption, may well have outweighed the risks involved. In other areas, at the frontiers of glass types, the adoption of the idea of recycling may have been accepted more gradually.

CONCLUSION

The widespread belief that glass in the Roman period was recycled lead to the general assumption that all glass could be recycled. While it is true that if a high enough temperature is reached (as would probably have happened in tank furnaces during primary glass manufacture), then most types of glass can be incorporated as cullet into a large batch. The inclusion of plant ash glass cullet in remelting a batch of natron glass, however, could have detrimental effects. In low quantities this may have been a negligible issue, but as the amount of plant ash glass used was increased, so too did the risk of devitrification increase. Natron glass, as such, could be recycled and reused. However, with early furnace technology, plant ash glass could not. It is most likely that 'old' glass or cullet would have been used as pieces or chunks of glass, since powdering would have created the nuclei necessary for crystallisation to occur. Yet, even in glass form, the composition of natron glass does not preclude it from devitrifying during recycling and subsequent manufacture. Mixing plant ash glass in with natron glass recycling would lead to increased crystal formation. The nature of this process would be determined partly by the composition of the melt and partly by the

temperature of the furnace. The horizontal heating chamber furnace used for glass blowing, which enabled the melt to stay hotter for longer, was invented in the 1st century BC. Prior to this glass workers may not have been able to maintain a high enough temperature to ensure a crystal free melt. This may explain why the discovery that glass could be completely remelted and therefore recycled did not occur until relatively late (c.1st century AD). The nature, amount and rate of crystal growth is dependent on a number of variables including melt composition, liquidus temperature, forming temperature, ΔT_{FL} , furnace temperature, crucible composition and cooling rate. Therefore, evidence of glass recycling in antiquity is testament to the skills and technical knowledge of the glass workers.

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Table 1: The percentages of the powdered glasses mixed together to form the new melts.

Table 2: Operating parameters of the ICP-OES.

Table 3: The results of the ICP-OES analyses of the samples, reported in wt%.

Table 4: The predicted compositions of the glasses mixed with plant ash glass in wt%.

Table 5: The liquidus temperatures and viscosity temperatures of the glasses determined using the liquidus temperature calculator and the viscosity calculator (<http://glassproperties.com/>). The modelled and actual SiO₂ values are also listed in wt%. The ΔT_{FL} and working range for each glass is also shown.

Figure 1: Sample RBS-D (75% plant ash glass, 25% natron glass) indicating the crystalline phases present. Left: plain polarised light, 5x magnification; right: plain polarised light, 20x magnification.

Figure 2: Thin sections of the glass indicating the increasing volume of crystals in relation to the increasing amount of plant ash glass in the melt. All glasses are shown in plain polarised light at 5x magnification. From left to right and from top to bottom (RBS-H 10% PA; RBS-C 25% PA; RBS-G 33% PA; RBS-F 50% PA; RBS-D 75% PA; RBS-PA 100% PA).

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