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Impact of recently discovered sodium calcium silicate solutions on the phase diagrams of relevance for glass-ceramics in the Na₂O-CaO-SiO₂ system

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ABSTRACT

Although phase relations in the Na₂O-CaO-SiO₂ system are vital to melting and thermal treatments in glass and glass-ceramics industries, the available data for thermodynamic modeling are mostly based on reports published in 1920s and 1950s. The present investigation verifies the formation of solid solutions of Na₂CaSiO₄ and Na₂Ca₂Si₂O₇ which have previously assumed to be stoichiometric compounds. The impact of these solid solutions on the features of the phase diagram were investigated using the equilibration-quenching-EDS/EPMA technique. The data were reported as liquidus projections and in isothermal sections within the temperature range of 1000 and 1400 °C. Ten primary phase fields were identified, namely SiO₂, Na₂Ca₃Si₆O₁₆, combeite, Na₄CaSi₃O₉ss, CaSiO₃, Na₂CaSiO₄ss, Na₂Ca₂Si₂O₇ss, Na₂Ca₆Si₄O₁₅, Ca₃Si₂O₇ and Ca₂SiO₄. In addition, some novel liquidus data and invariants points were examined in more detail. The fundamental data obtained can be employed for the thermodynamic reassessment of the Na₂O-CaO-SiO₂ system. The present study also discusses the findings and their impact on melting and annealing processes during the manufacture of glass and glass-ceramics.

1. Introduction

Soda-lime-silica is the most important glass family in daily life because 90 % of the manufactured conventional glasses in the world belong to the Na₂O-CaO-SiO₂ system [1]. The Na₂O-CaO-SiO₂ glass system is also vital in many medical applications, including several compositions used for tissue engineering and drug delivery [2,3]. Understanding the behavior of the Na₂O-CaO-SiO₂ system, e.g., the phase relations among those present at high temperatures, is therefore crucial for manufacturing these glasses and ceramics. One tool to help manufacturers in the essential task of successfully predicting the melting range of a particular mixture and the crystalline phases formed during any subsequent heat treatment is the detailed understanding of the phase diagram Na₂O-CaO-SiO. However, the data used to construct this diagram is based on the classic papers by Kracek [4] and Morey and Bowen [5] in 1925 and further developed by Segnit [6], as resumed in the book of Phase Diagram for Ceramist [7]. For the metallurgical industries, the system is vital for slags as Na₂O could significantly decrease their melting point and increase a slag's capacity to remove impurities [8]. For the incineration technology to process the Municipal Solid Waste (MSW), the system is important as the slag, accounted for 85 % by weight formed in the bottom of the reactors as the solid residue, contains considerable amounts of Na₂O, CaO and SiO₂ [9,10]

Morey and Bowen [5] experimentally investigated the phase relations of liquid with solid phases in the Na₂O-CaO-SiO₂ system by using the melting-quenching technique. The samples were prepared from the mixture of CaCO₃, Na₂CO₃ and SiO₂ heated using platinum crucible to form a glass which was used for the phase equilibria measurement. The glass composition was analyzed before the equilibration to measure phase relations. The quenched phases were examined by using optical microscope. They reported SiO₂, CaSiO₃, Na₂SiO₃, Na₂Si₂O₅, Na₂Ca₂. Si₃O₉, Na₈Ca₃Si₅O₁₇, Na₄CaSi₃O₉, and Na₂Ca₃Si₆O₁₆ primary phase fields. Then, Segnit [6] used the same experimental method used by Morey and Bowen [5] extended the study to the area containing less

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Table 1

Chemicals used in the present Investigation.

No	Chemical	purity, wt-%	Supplier
1	SiO ₂	99.9	Umicore
2	CaO	99.9	Sigma Aldrich
3	Na ₂ CO ₃	99.9	Sigma Aldrich
4	Na ₂ SiO ₃	96	Alfa Aesar

 SiO_2 and reported a new compound of $Na_8Ca_3Si_5O_{17}$ and measured liquidus data in the new primary phase fields $Na_8Ca_3Si_5O_{17}$, Ca_2SiO_4 , $Ca_2Si_2O_7$, Na_2CaSiO_4 , and $Na_2Ca_2Si_2O_7$. He [6] experienced difficulty at high temperatures due to soda volatilization. Later, Sahid and Glasser [11] revised the diagram after finding new crystalline phases of $Na_2CaSi_5O_{12}$ and $Na_6Si_8O_{19}$ and measured their liquidus points employing the same methods as Morey and Bowen [5].

However, the chemical composition measurements of the glasses reported by Morey and Bowen [5], Segnit [7], Sahid and Glasser [11] were undertaken before the re-melting in the equilibration process during which liquidus of the mixtures was measured. The phases in the quenched samples were examined using optical microscopy. As Na₂O is a volatile component, losing Na₂O to the atmosphere during the melting is likely to change the concentration of Na2O compared to the initial mixture. With regard to this limitation, post-equilibrium chemical analysis of the sample by employing the Equilibration-Quenching-EPMA method may be the solution [12]. Zhang et al. [13] also employed this method, although they quenched their samples in a nitrogen atmosphere that gives a slow quenching rate. As indicated in their results, the quenched liquid phases were not homogenous because they turned into microcrystalline phases during the quenching. This change proves that the liquid properties stable at high temperatures cannot be preserved to room temperature as the melt's composition might alter during the quenching process. Therefore, it is difficult to indicate the composition of the phases formed and phase relations from their results [13].

The existence of two solid solutions in the ternary system was reported by Moir and Glasser [14] when they reexamined the stability of combeite (Na₂Ca₂Si₃O₉) and Na₄CaSi₃O₉ compounds characterized by using XRD analysis for the samples synthesized using Morey and Bowen's techniques [5]. They showed that combeite was not a stoichiometric compound of Na2Ca2Si3O9, neither Na4CaSi3O9. These compounds form solid solutions existing in the pseudo-binary Na2SiO3-CaSiO3 system. Combeite has a compositional range of Na₂SiO₃ between 28.5 and 53.5 mol-%, whereas the Na₄CaSi₃O₉ has a much smaller range. They also reported solid solution of β-CaSiO₃ that can dissolve up to 2 mol-% Na2SiO3. Fedorov and Brodkina [15] outlined a solid solution of β -Ca₂SiO₄ in the pseudo-binary Ca₂SiO₄ and Na2CaSiO4. B-Ca2SiO4 was found to dissolve up to 10 wt-% Na2CaSiO4 at 1300 °C at which the sample was annealed and then characterized by XRD method. Toropov and Arakelyan [16] identified two new compounds in the Ca₂SiO₄ and Na₂CaSiO₄ system, hexagonal Na₄Ca₈Si₅O₂₀ and Na₄Ca₄Si₃O₁₂. However, according to Fedorov and Brodkina [15], Na₄Ca₈Si₅O₂₀ is actually a β-Ca₂SiO₄ solid solution and Na₄Ca₄Si₃O₁₂ does not exist.

In terms of thermodynamic modeling, Thermo-Calc [18] was employed by L. Zhang et al. [17] to assess the Na₂O-CaO-SiO₂ system. The sub-lattice model was used to describe the Gibbs energy of liquid. Later, Z. Zhang et al. [19] assessed the system using a modified associated species model to evaluate the Gibbs energy of the liquid phase and employing FactSage using FToxid database [20] for the calculation. Some disagreements were obvious between the results in these two works by Z. Zhang et al. and L. Zhang et al. For example, in the diagram produced by L. Zhang et al. [17], the pseudo-binary Na₂SiO₃-CaSiO₃ section was much more complicated than the one presented by Z. Zhang et al. [19]. It is because the computed quasi-binary diagram by L. Zhang et al. [17] did not indicate the stability of the Ca₂SiO₄, Na₂Si₆O₇, and Na₂Ca₂Si₃O₉ compounds. However, these compounds were stable in the computed quasi-binary diagram by Z. Zhang et al. [19].

Kahlenberg et al. [21] recently reported the Na₂Ca₆Si₄O₁₅ compound. Its liquidus data have not been available in the literature. Therefore, the inclusion of this compound in the thermodynamic assessment of the Na₂O-CaO-SiO₂ system will affect the computed phase diagrams. They [21] synthesized the compound by using Na₂CO₃-Ca-CO₃-SiO₂ placed in a platinum container and heated 1300 °C and then quenched. The sample was characterized by using EPMA to measure chemical compositions and by employing X-ray diffractometer to collect crystallographic data. The existence of some compounds, such as Na₈Ca₃Si₅O₁₇, Na₂Ca₃Si₂O₈ and Na₂Ca₃Si₅O₁₂, was questioned [22] as they cannot be reproduced experimentally by using the crystallization approach and solid-state syntheses.

It is obvious that there is a lack of reliable experimental liquidus data for mixtures with higher Na₂O contents. The omission of liquidus data for the newly reported crystals is evident. Disagreement among the authors [17,19] is also apparent. To conclude the above analysis, to better understand the formation of suggested additional crystalline phases and their liquidus, experimental reinvestigation of the system Na₂O-CaO-SiO₂ is required. In the present study, the Equilibration-Quenching-EPMA/EDS method was employed to study the Na₂O-CaO-SiO₂ system over a wide composition range of SiO₂ from 33.33 to 84.2 mol-% and Na2O up to 47.5 mol-% between 1000 and 1400 °C. The primary phase fields were explored to derive the saturation boundary lines of the compounds. Furthermore, liquidus with double saturation was also studied to obtain the univariant lines, which are the boundary lines between two adjacent primary phase fields. Comparison of the data obtained in the present work to previous research, as well as the computed phase diagram was undertaken. The results are discussed in relation to the manufacturing of the glass.

2. Methods

When investigating the equilibrium properties of phases occurring at high temperatures, the phases must be preserved by rapid cooling in a cold medium. Then, the composition and homogeneity of the phases can be examined under a microanalyzer. The liquid must be preserved as a glassy phase, and also the solid phase composition must be maintained. In the present investigation, the sample preparation challenges, were met by utilizing the Equilibration-Quenching-EPMA/EDS technique.

2.1. Sample preparation

The high purity chemicals presented in Table 1 were used for the preparation of the initial mixtures. The chemicals were weighed and mixed carefully using an agate mortar and pestle, then pressed to form a 0.15 g pellet. It was important to know the degree of Na₂O volatilization during the equilibration process; thus, the excess amount of Na₂O in the samples was specified accurately to obtain the targeted phase relations. Failing to take into account the volatilization would result in unwanted phase equilibria as the evaporation changes the location of the tie lines and the proportions between equilibrium phases.

Na₂O concentration in the mixture was adjusted to form a liquid saturated with one or several crystalline phases. Otherwise, if the mixture formed upon quenching only amorphous phase(s), the liquidus boundaries would not be revealed. For this purpose, the diagram computed by FactSage using FToxid database [20] and MTDATA [23] using MTOX database [24] were used as the initial guidance for the mixtures.

2.2. Equilibration process

The pellets were placed in a platinum container and held by a platinum wire in the center of the hot zone of the reaction tube with 30 mm inner diameter, made from impervious recrystallized alumina, inside a furnace (Nabertherm RHTV 120-150/18, heated with MoSi₂ heating I. Santoso et al.



Fig. 1. Furnace arrangement.



Fig. 2. Sample container design.

Table 2

Comparison between EDS analysis results and stoichiometric values of the formed crystalline compounds.

Analysis	Na ₂ Ca ₆ Si ₄ O ₁₅ composition (mol-%)			Na ₂ Ca ₃ Si ₆ O ₁₆ composition (mol-%)			
	Na ₂ O	CaO	SiO ₂	Na ₂ O	CaO	SiO ₂ 60.0	
True stoichiometric value EDS	9.1 9.2	54.6 54.8	36.4 36.0	10.0 10.2	30.0 29.5	60.0 60.3	

Table 3

Comparison between EDS and EPMA analysis results of the solid and liquid phases.

Apolizio	Solid cor	mposition ((mol-%)	Liquid composition (mol-%)			
Allalysis	Na ₂ O	CaO	SiO_2	Na ₂ O	CaO	SiO_2	
EPMA EDS	26.0 26.5	40.2 39.7	33.8 33.8	27.8 28.2	38.7 38.2	33.6 33.6	

element, Germany) to equilibrate the samples. The arrangement of the furnace and the design of the Pt crucible can be seen in Figs. 1 and 2, respectively. The top of the reaction tube was equipped with two holes

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for the thermocouple sheath and one for a mini tube to protect the wire. A calibrated S-type thermocouple of Pt/Pt-10 wt-% Rh (Johnson-Matthey, UK) was placed close to the samples and connected to a multimeter (Keithley, USA), which was linked to a computer program (NI LabView) to continuously measure and document the temperature of the samples every 5 s. Temperature uncertainty of ± 2 °C was achieved. The bottom of the reaction tube was open to the air.

2.3. Equilibrium confirmation

Equilibrium attainment can be ensured from three different criteria: time, the direction of approach, and homogeneity. An equilibrium has been reached if samples heated at different equilibration times but at the same temperature locate on the same liquidus line. Therefore, samples in the present investigation were equilibrated at different holding times, and the results showed that the sample could reach equilibrium after 1 h or 2 days depending on temperature and the specific type of equilibria. The second criterion means that the same equilibrium state must be reachable either by heating the sample directly to the targeted temperature or by pre-melting, i.e., overheating the sample and then cooling it down to the targeted temperature. Also, the criterion is satisfied if different compounds used as the initial mixture can produce the same equilibrium condition. In the present investigation, mixtures of Na₂CO₃-CaO-SiO₂ or Na₂SiO₃-CaO-SiO₂ were employed (used as the main mixtures) to ensure that the solid phase in the final state was formed by chemical reactions, not from the unreacted oxides in the initial mixture.

The equilibrium state can be attained as well from the direction of the right side with lower Na₂O content and from the left side with higher Na₂O content. For example, the equilibrium state of Na₂Ca₂Si₂O₇ solid solution (Na₂Ca₂Si₂O₇ss) can be reached from the Na₂Ca₂Si₃O₉ solid solution (combeite) or the Na₂CaSiO₄ solid solution (Na₂CaSiO₄ss). From the thermodynamic point of view, if the equilibria of liquid-Na₂Ca₂Si₂O₇ss exist, the equilibria of liquid-Na₂Ca₂Si₂O₇ss-Na₂CaSiO₄ss and equilibria of liquid-Na₂Ca₂Si₂O₇ss-combeite must exist as well. According to the third criterion, no concentration gradient in any phase is allowed. Hence, the liquidus microanalyses were undertaken at multiple locations to ensure an equal concentration throughout the phases.

2.4. Quenching

After the equilibrium was achieved, the sample was dropped to cooled water by pulling upward the platinum wire that holds the sample. The quenched samples were then dried immediately using hot flash air and mounted in epoxy resin. The sample was polished by using a dry technique, carbon coated and sent for chemical analysis. There was no difficulty to obtain samples which contained glassy phase.

2.5. Phase examination

The samples were analyzed using EPMA (JXA-8530 F Plus Hyper Probe, Jeol, Japan) at Center for Material Analysis, University of Oulu and EDS (ThermoFisher Scientific UltraDry, USA) installed in SEM (Tescan Mira3, Tescan, Czech Republic) at Aalto University. Optimized instrument parameters were needed because of matrix modification resulted from the sodium ion migration in the sample during measurement. For the EPMA measurement 15 kV, 15 µm and 10 s of accelerating voltage, beam diameter, measuring time, respectively, were selected to measure the glass/amorphous phases. For the EDS measurement, area analysis of 100-200 µm², 15 kV accelerating voltage and 420 nA current were employed. Tugtupite (Na4BeAlSi4O12Cl), calcite (CaCO3) and quartz (SiO₂) minerals were used as Na, Ca and Si standards, respectively. ZAF correction was used for EPMA and EDS analyses. A comparison of results between EDS measured values and true stoichiometric values are presented in Table 2. Table 3 shows a comparison of the result obtained with EDS and EPMA analyses of the Na2CaSiO4ss compound and its amorphous phase representing the liquid phase at the high

Table 4

Phase compositions measured by EDS analysis.

	<u> </u>			Composition (mol-%))	
Sample #	Temperature	Equilibration time (h)	Phases	Na ₂ O	CaO	SiO ₂
N/00 1 4/	1000		Liquid	15.7	11.7	72.6
NCS-146	1000	55.5	Na ₂ Ca ₃ Si ₆ O ₁₆	10.2	29.5	60.3
NCS-147	1000	55.5	Liquid	17.5	11.9	70.6
			$Na_2Ca_3Sl_6O_{16}$	10.2	29.8 12.6	60 67 4
NCS-148	1000	55.5	Na ₂ Ca ₃ Si ₆ O ₁₆	10.4	29.7	59.9
			Combeite	16.8	33.3	49.9
NCS-150	1000	55.5	Liquid	24	9.9	66
1100 100	1000		Combeite	19.1	31	49.9
NCS-151	1000	55.5	Liquid	28.2	7.5 28 3	64.3 49.6
	1000		Liquid	40.2	2.9	57
NCS-152	1000	43	Na4CaSi3O9ss	32.1	17.1	50.7
			Liquid	36	4.3	59.7
NCS-155	1000	43	Na ₄ CaSi ₃ O ₉ ss Combeite	32.3	17.5	50.2
			Liquid	33.9	5	49.9 61.1
NCS-157	1000	43	Combeite	24.2	26.2	49.6
NCS-158	1000	43	Liquid	26.1	8.6	65.3
			Combeite	20.7	29.5	49.8
NCS-159	1000	43	Liquid Combeite	31./ 23.4	5.8 27	62.5 49.7
N/00.1/0	1000	40	Liquid	17.8	5.9	76.3
NCS-160	1000	43	SiO ₂	0	0	100
			Liquid	36.8	4.2	59.1
NCS-162	1000	43	Na ₄ CaSi ₃ O ₉ ss	32.6	17.3 25 5	50.1
			Liquid	15.4	23.3 10.1	49.9 74.5
NCS-163	1000	43	SiO ₂	0	0	100
NCS-338	1000	30	Liquid	21.6	11.3	67.1
1100 000	1000		Combeite	17.7	32.4	49.9
NCS-341	1000	30	Liquid Combeite	20.4 17	33	67.7 50
N/00 100	1100		Liquid	40.6	15.8	43.6
NCS-183	1100	6	Na ₂ CaSiO ₄ ss	30.7	35.5	33.8
		_	Liquid	37.5	18.6	43.9
NCS-190	1100	7	$Na_2Ca_2Si_2O_7ss$ No. CoSiO, co	20.3	40.2	39.5
			Liquid	37.1	37 17.5	33.7 45.4
NCS-192	1100	7	Na2Ca2Si2O7ss	20	40.4	39.6
			Liquid	36.7	17.4	45.9
NCS-193	1100	7	Na ₂ Ca ₂ Si ₂ O ₇ ss	20.3	40.2	39.5
			Liquid	32.3 36.1	7.6	49.7 56.3
NCS-196	1100	8	Na ₄ CaSi ₃ O ₉ ss	32.1	18	49.9
			Combeite	24.4	25.8	49.8
NCS-199	1100	8	Liquid	29.1	9.4	61.6
			Liquid	21.3 22.1	29 14.8	49.7 63.1
NCS-200	1100	24	Combeite	18.4	31.8	49.8
NCS-204	1100	24	Liquid	19.8	16.2	64
100 201	1100	21	Combeite	16.9	33.2	49.9
NCS-205	1100	24	Liquid CaSiO	17.4	17.9	64.7 49 5
103-203	1100	27	Combeite	15.1	34.9	49.5 50
NCS-207	1100	24	Liquid	16.3	16.4	67.4
NG3-207	1100	27	CaSiO ₃	0	50.1	49.9
NCS-209	1100	24	Liquid	14.6	15.3	70.2
			Liquid	12.1	14.2	49.9 73.7
NCS-214	1100	24	SiO ₂	0	0	100
			CaSiO ₃	0	49.9	50.1
NCS-216	1100	24	Liquid	14.6	8.8	76.7
			5102 Liquid	0 17	4	100 79
NCS-217	1100	24	SiO ₂	0	0	100
NCS-8	1200	75	Liquid	13.1	6.4	80.6
1.00 0	1200		SiO ₂	0	0	100
NCS-10	1200	75	Liquid	8.3 0	18.7	73
1100-10	1200	/5	CaSiO ₃	0	50.3	49.7
NCS 12	1200	75	Liquid	10.5	20.8	68.7
1103-12	1200	15	CaSiO ₃	0	50.2	49.8

(continued on next page)

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Table 4 (continued)

Sample #	Temperature	Equilibration time (b)	Phases	Composition ((mol-%)		
Sample #	remperature	Equilibration time (ii)	1 110505	Na ₂ O	CaO	SiO_2	
NCS-13	1200	44	Liquid	10.7	20.7	68.6	
			CaSiO ₃ Liquid	0	50.3 21.8	49.7 66 9	
NCS-14	1200	44	CaSiO ₃	0	50.4	49.6	
NCS-15	1200	44	Liquid	13.3	23.4	63.3	
1100 10	1200		CaSiO ₃ Liquid	0	50.3 25 4	49.7 61	
NCS-16	1200	44	CaSiO ₃	0	50.2	49.8	
NCS-19	1200	72	Liquid	19	22.8	58.1	
			Combeite	16.6	33.4	50	
NCS-28	1200	68	Combeite	19.8	30.5	49.7	
NCS-81	1200	6	Liquid	34	22.9	43.1	
			Na ₂ CaSiO ₄ ss Liquid	27	39.3 17	33.6 52.1	
NCS-85	1200	6	Combeite	22.1	28.2	49.7	
NCS-99	1200	6	Liquid	30	25.3	44.7	
			Na ₂ Ca ₂ Si ₂ O ₇ ss Liquid	18.6 31.2	41.6 25.9	39.8 42 9	
NCS-102	1200	6	Na ₂ Ca ₂ Si ₂ O ₇ ss	20.1	40.7	39.1	
			Na ₂ CaSiO ₄ ss	26.5	39.7	33.8	
NCS-107	1200	46	Liquid Na-Ca-Si-O-ss	26.7 17.3	27.2 43.1	46.1 39.6	
103-107	1200	10	Combeite	21.5	28.9	49.6	
NCS-115	1200	22	Liquid	20.4	21	58.6	
			Combeite	16.7 20.4	33.6 21	49.7 58.6	
NCS-116	1200	22	Combeite	16.8	33.5	49.8	
NCS-117	1200	22	Liquid	23.1	19.3	57.6	
1100 117	1200		Combeite	18.1	32.3	49.7	
NCS-120	1200	22	Combeite	13.1	36.3	49.9	
			CaSiO ₃	0	50.5	49.5	
NCS-166	1200	21	Liquid Combeite	29.6	22.6	47.9	
	4000		Liquid	30.6	19.6	49.0	
NCS-167	1200	21	Combeite	23.4	27	49.6	
NCS-285	1200	1	Liquid	47.5	12.7	39.8	
	4000		Liquid	42.5	15.7	41.8	
NCS-290	1200	2	Na ₂ CaSiO ₄ ss	31.1	35.6	33.3	
NCS-294	1200	2	Liquid	40.2	18	41.8	
1000 0.45	1000	-	Liquid	37.2	20.2	42.7	
NCS-345	1200	5	Na ₂ CaSiO ₄ ss	28.9	37.6	33.5	
NCS 26	1200	24	Liquid	5.4	24.1	70.5	
NG3-30	1300	24	CaSiO ₃	0	50.2	49.8	
NCS-40	1300	24	Liquid	9.7	29.7	60.6	
1105-40	1300	27	CaSiO ₃	0	50.3	49.7	
NCS-42	1300	24	CaSiO ₃	0	20.9	49.5	
NCS-43	1300	24	Liquid	7.4	48.1	44.5	
	1000		Ca ₂ SiO ₄ ss	0.3	66.4 42.4	33.3	
NCS-45	1300	24	CaSiO ₃	9.7 0	50.2	47.9	
NCS-48	1300	24	Liquid	8.4	47.4	44.2	
			Na ₂ Ca ₆ Si ₄ O ₁₅	7.7	56.2	36.2	
NCS-50	1300	24	Na ₂ Ca ₆ Si ₄ O ₁₅	8.4	55.7	35.8	
NCS-137	1300	4 5	Liquid	16	39.2	44.8	
	1000		Na ₂ Ca ₆ Si ₄ O ₁₅	8.7	55.1	36.2	
NCS-170	1300	5	SiO ₂	0	9.1 0	80.2 100	
NCS-171	1300	5	Liquid	8.9	13.7	77.3	
		-	SiO ₂ Liquid	0	0	100	
NCS-172	1300	5	CaSiO ₃	0	50.8 50.5	49.5	
NCS-179	1300	13	Liquid	11.7	43.3	45	
	1000		Na ₂ Ca ₆ Si ₄ O ₁₅	8.6	55.1	36.4	
NCS-300	1300	12	CaSiO ₃	0	50.2	45.4 49.9	
			Ca ₃ Si ₂ O ₇	0	59.9	40.1	
NCS-315	1300	1	Liquid	30.1	27.8	42.1	

(continued on next page)

Table 4 (continued)

0 1 "			DI.	Composition (Composition (mol-%)	
Sample #	Temperature	Equilibration time (h)	Phases	Na ₂ O	CaO	SiO ₂
			Na ₂ CaSiO ₄ ss	25.8	41	33.2
			Liquid	26.4	31.8	41.8
NCS-350	1300	1	Na ₂ Ca ₆ Si ₄ O ₁₅	9.2	54.8	36
			Na ₂ CaSiO ₄ ss	24.2	42.6	33.2
NCS 262 a	1200	1	Liquid	37.1	21	41.9
NG3-302-a	1300	1	Na ₂ CaSiO ₄ ss	29.1	37.6	33.3
NCS 362	1300	1	Liquid	33.7	24.8	41.5
NG3-302	1300	1	Na2CaSiO4ss	27.9	38.7	33.4
NCS-362 NCS-55 NCS-58 NCS-63	1400	20.5	Liquid	8.1	7.7	84.2
1163-33	1400	20.3	SiO ₂	0	0	100
NCS 58	1400	20.5	Liquid	5.4	19.1	75.5
1103-30	1400	20.3	SiO ₂	0	0	100
NCS 63	1400	28	Liquid	2.7	33.9	63.4
NC3-03	1400	20	CaSiO ₃	0	50.4	49.6
NCC 6E	1400	20	Liquid	6.2	36.8	0 100 33.9 63.4 50.4 49.6 36.8 57 50.4 49.6
NCS-65	1400	28	CaSiO ₃	0	50.4	49.6
NCS 60	1400	28	Liquid	9.7	47.3	43
NCS-65 NCS-69	1400	20	Ca ₂ SiO ₄ ss	2.1	64.3	33.6
NCS 73	1400	4.5	Liquid	16.8	41	42.2
NCS-63 NCS-65 NCS-69 NCS-73 NCS-74	1400	4.5	Na ₂ Ca ₆ Si ₄ O ₁₅	8.8	55	36.2
NCS 74	1400	4.5	Liquid	14.2	43.3	42.5
1103-74	1400	4.5	Na ₂ Ca ₆ Si ₄ O ₁₅	8.5	55.5	36
			Liquid	3.6	52.5	43.9
NCS-305	1400	6	CaSiO ₃	0	50.3	49.7
			$Ca_3Si_2O_7$	0	60	40
NCS 328	1400	1	Liquid	25.3	33.8	40.8
1103-520	1400	1	Na ₂ CaSiO ₄ ss	23.4	43.4	33.2
			Liquid	22.5	36.8	40.8
NCS-358	1400	1	Na ₂ Ca ₆ Si ₄ O ₁₅	9.2	54.5	36.4
			Na ₂ CaSiO ₄ ss	22.5	44.3	33.3



Fig. 3. Method employed in the present investigation.

temperature equilibrium condition.

It can be seen in Table 2 that the EDS results are in very close agreement with the theoretical values. They are also in good agreement with the EPMA results, as indicated in Table 3. The maximum uncertainties obtained were less than 1 mol-%. In the present study, EDS analysis results of liquid phase are presented together with the results of solid phase analysis, in Table 4. Therefore, the analysis accuracy of each amorphous phase, i.e., liquid phase at the high temperature, also can be checked from the results of the respective solid phase. The method is summarized in Fig. 3.

2.6. Computational phase diagram

The computed phase diagrams presented were calculated by FactSage 7.3. using FToxid database [20] and MTDATA [23] using MTOX database, version 8.2 [24]. The obtained phase diagrams were then compared with the experimental data.

3. Result and discussion

The compositional analyses of the amorphous (liquid) and solid phases using EDS are reported in Table 4. In the present study, 10 primary phase fields were obtained since altogether 10 solid compounds were found to exist in equilibrium with the liquid phase and verified in the quenched samples. These were the phase fields of SiO₂, Na₂Ca₃. Si₆O₁₆, combeite (Na₂Ca₂Si₃O₉), Na₄CaSi₃O₉ss, CaSiO₃, Na₂CaSiO₄ss, Na₂Ca₂Si₂O₇ss, Na₂Ca₆Si₄O₁₅, Ca₃Si₂O₇ and Ca₂SiO₄. The micrographs of a liquid saturated with a single and two types of crystals are presented in Figs. 4 and 5 respectively. The detailed relation among the phases obtained is presented as isothermal sections at 1000, 1100, 1200, 1300 and 1400 °C in Figs. 6–10, respectively. In the isothermal sections, this liquid is represented by tie lines connecting the liquid with the solid compound. The liquid phase is graphically presented as liquidus projections in Fig. 11. The solid phases connected by Alkemade lines are reported in Fig. 12. Each primary phase field intersects with the adjacent



Fig. 4. Backscattered electron micrographs of amorphous phase, i.e. liquid in equilibrium with A. SiO₂; B. Na₂Ca₃Si₆O₁₆; C. Combeite; D. Na₄CaSi₃O₉ss; E. CaSiO₃; F. Na₂CaSiO₄ss; G. Na₂Ca₂Si₂O₇ss; H. Na₂Ca₆Si₄O₁₅; I. Ca₂SiO₄.

fields along the univariant lines drawn as a solid black line in Fig. 11. The microstructures are represented by the amorphous phase in equilibrium with two solid phases in Fig. 5. In the isothermal sections, this liquid is depicted by the tie triangle connecting the liquid with the two solid compounds.

Altogether, 18 different phase assemblages of the quenched samples were obtained in the present study. These assemblages are collected in Table 5: Liquid + SiO₂, liquid + Na₂Ca₃Si₆O₁₆, liquid + combeite, liquid + Na₄CaSi₃O₉ss, liquid + CaSiO₃, liquid + Na₂Ca₂SiO₄ss, liquid + Na₂Ca₂Si₂O₇ss, liquid + Na₂Ca₆Si₄O₁₅, liquid + Ca₂SiO₄ss, liquid + SiO₂+CaSiO₃, liquid + combeite + Na₂Ca₃Si₆O₁₆, liquid + combeite + Na₄CaSi₃O₉ss, liquid + combeite + CaSiO₃, liquid + combeite + Na₂Ca₂Si₂O₇ss, Liquid + Na₂CaSiO₄ss + Na₂Ca₆Si₄O₁₅, liquid + Na₂Ca₂Si₂O₇ss + Na₄CaSi₃O₉ss, liquid + Na₄Ca₂Si₄O₇s, and liquid + CaSiO₃ + Ca₃Si₄O₇.

Comparisons between the experimental data and the calculated phase diagrams by FToxid database [20] are presented in Figs. 13 and 14 as $CaSiO_3$ -Na₂SiO₃ and Ca_2SiO_4 -Na₄SiO₄ pseudo binary diagrams, respectively. Figs. 15–17 show comparisons between the computed diagrams by FToxid [20] and MTOX databases [24] and the data at 1000, 1200 and 1400 °C, respectively.

3.1. Behavior of Na₂O

The study of the Na₂O evaporation was similar to that used in an earlier study of K₂O [25]. Fig. 18 shows how the tie lines move due to the evaporation of Na₂O. The remaining Na₂O content in the equilibrium phase at a specific temperature depended on the equilibration time and the concentration of Na₂O in the initial mixture. If Na₂O evaporates gradually, the composition will also change gradually following the

evaporation line toward the CaO-SiO₂ side, drawn as a dashed line. As shown in Fig. 18, Na₂O concentration in sample NCS-81 decreased from 42.5 mol-% to 31 mol-% after being equilibrated for just 6 h at 1200 °C. However, sample NCS-13 with Na₂O concentration of 10 mol-% in the initial mixture decreased to 7.5 mol-% after 44 h equilibration. Therefore, to obtain a particular final equilibrium assemblage, the specific concentration in the initial mixture must be carefully adjusted.

3.2. Primary phase field of SiO₂

The liquidus point data in SiO2 primary phase field at 1000 °C are in good agreement with the FToxid database [20] and have slightly lower Na₂O solubility than that calculated with the MTOX database [24] (Fig. 15). The data agree with those reported by Santoso and Taskinen [12], who measured the liquidus data at SiO₂ saturation in the binary Na2O-SiO2 system. MTDATA and its MTOX database [24] used the data by Kracek [4] and Morey and Bowen [5] in its assessment where the calculated liquidus line will have higher Na₂O concentration. Thus, the measured liquidus line should be lower in Na2O content compared to the MTOX database simulation due to some Na2O evaporation in the experiments by Kracek [4] and Morey and Bowen [5]. The present study reports liquid data in double saturation of SiO₂ and CaSiO₃ (Fig. 5A), thus providing data to locate the intersection between the SiO₂ and CaSiO₃ primary phase fields. The locations of intersection between SiO₂ and CaSiO₃ in the ternary diagram has a higher CaO content than calculated by the FToxid database [20], as indicated in Figs. 16 and 17.

3.3. Primary phase field of CaSiO₃

Between 1200 and 1400 °C, liquidus point data at CaSiO₃ saturation



Fig. 5. Backscattered electron micrographs of amorphous phase (liquid) in equilibrium with A. $SiO_2 + CaSiO_3$; B. $Na_2Ca_3Si_6O_{16} + Combeite$; C. Combeite + $Na_4CaSi_3O_3Ss$; D. Combeite + $CaSiO_3$; E. Na_2CaSiO_4ss + $Na_2Ca_6Si_4O_{15}$; F. $Na_2Ca_6Si_4O_{15}$ + Na_2CaSiO_4ss ; G. $Na_2Ca_2Si_2O_7ss$ + $Na_4CaSi_3O_9ss$; H. Na_2CaSiO_4ss + $Na_2Ca_2Si_2O_7ss$; I. $CaSiO_3 + Ca_3Si_2O_7$.



Fig. 6. Isothermal section of the Na₂O-CaO-SiO₂ system at 1000 °C.

are lower in Na₂O content than the liquidus line predicted by FToxid [20] and MTOX databases [24] (Figs. 16 and 17). This difference can be explained by the fact that these two softwares are, again, based on Kracek [4] and Morey and Bowen [5] data. Evaporation of Na₂O should have shifted their data systematically away from the Na₂O corner. According to Moir and Glasser [14], CaSiO₃ can dissolve Na₂SiO₃ up to 5 mol-%. However, the present investigation proved that solid CaSiO₃ obtained at 1200 and 1300 °C contains only a negligible amount of Na₂O. Liquid compositions at double saturation of CaSiO₃ and Ca₃Si₂O₇



Fig. 7. Isothermal section of the Na₂O-CaO-SiO₂ system at 1100 °C.

(Fig. 5I) were obtained at 1300 and 1400 °C, meaning that the intersection location between primary phase fields of $CaSiO_3$ and $Ca_3Si_2O_7$ is known in the ternary diagram.

3.4. Primary phase field of Ca₃Si₂O₇

The primary phase field of Ca₃Si₂O₇ obtained in the present study is very narrow, as shown in Fig. 11. The width of the liquidus contours is relatively constant at low and high Na₂O concentrations, which agrees



Fig. 8. Isothermal section of the Na2O-CaO-SiO2 system at 1200 °C.



Fig. 9. Isothermal section of the Na₂O-CaO-SiO₂ system at 1300 °C.

with the Segnit [6] report and the values computed with the MTOX database [24]. This kind of feature of the $Ca_3Si_2O_7$ primary phase field was also found to be typical to the $Ca_3Si_2O_7$ primary phase field in the Ka₂O-CaO-SiO₂ system [25]. It differs, however, from the line computed with the FToxid data [20]. The computed field becomes much wider toward the Na₂O-SiO₂ direction. Nevertheless, as shown in Fig. 17, the location of the intersection between the Ca₂SiO₃ and Ca₃Si₂O₇ primary phase fields is in good agreement with the predictions of FToxid [20] and MTOX databases [24].

3.5. Primary phase field of Ca₂SiO₄

In the present study, the primary phase field Ca_2SiO_4 was observed at 1300 and 1400 °C. At 1400 °C, Ca_2SiO_4 dissolved 3.1 mol-% Na_4SiO_4 and appears in the pseudo-binary Ca_2SiO_4 - Na_4SiO_4 section (Fig. 14). It means that Ca_2SiO_4 formed a solid solution extending to the direction of Na_2CaSiO_4ss , and the maximum solubility of Na_4SiO_4 can be greater than 3.1 mol-% Na_4SiO_4 . This is consistent with the results of Fedorov and Brodkina [15]. The primary phase field of Ca_2SiO_4 intersects with the $Na_2Ca_6Si_4O_{15}$ field at 1300 and 1400 °C, as well as the $Ca_3Si_2O_7$ field, whereas FToxid [20] and MTOX databases [24] predict that at



Fig. 10. Isothermal section of the Na₂O-CaO-SiO₂ system at 1400 °C.



Fig. 11. Liquidus projection and univariant lines (black) of the Na_2O -CaO-SiO₂ system.

1300 and 1400 $^{\circ}$ C, Ca₂SiO₄ directly crosses the CaO primary phase field. Liquidus at Ca₂SiO₄ saturation obtained in the present investigation is in good agreement with the MTOX database [24] computation.

3.6. Primary phase field of Na₂Ca₆Si₄O₁₅

Na₂Ca₆Si₄O₁₅ crystals were formed in the sample equilibrated at 1300 and 1400 °C, thus confirming the observations by Kohlenberg [21]. In addition, the present study successfully measured novel liquidus data at Na₂Ca₆Si₄O₁₅ saturation. The intersection between the Na₂Ca₆Si₄O₁₅ and Na₂CaSiO₄ss primary phase fields between 1300 and 1400 °C was finally confirmed since the liquid phase in the double saturation of Na₂Ca₆Si₄O₁₅ and Na₂CaSiO₄ss was verified (Fig. 5F). This compound, Na₂Ca₆Si₄O₁₅, is not present in FToxid database [20] and MTOX database [24].



Fig. 12. Invariants points and the Alkemade lines in the $\ensuremath{\text{Na}_2\text{O-CaO-SiO}_2}$ system.

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Phase assemblages and their microstructures in the present investigation.

No	Phase assemblages	Microstructure
1	$Liquid + SiO_2$	Fig. 4A
2	$Liquid + Na_2Ca_3Si_6O_{16}$	Fig. 4B
3	Liquid + combeite	Fig. 4C
4	$Liquid + Na_4 CaSi_3O_9 ss$	Fig. 4D
5	$Liquid + CaSiO_3$	Fig. 4E
6	$Liquid + Na_2CaSiO_4ss$	Fig. 4F
7	$Liquid + Na_2Ca_2Si_2O_7ss$	Fig. 4G
8	$Liquid + Na_2Ca_6Si_4O_{15}$	Fig. 4H
9	$Liquid + Ca_2SiO_4$	Fig. 4I
10	$Liquid + SiO_2 + CaSiO_3$	Fig. 5A
11	$Liquid + combeite + Na_2Ca_3Si_6O_{16}$	Fig. 5B
12	$Liquid + combeite + Na_4 CaSi_3O_9 ss$	Fig. 5C
13	$Liquid + combeite + CaSiO_3$	Fig. 5D
14	$Liquid + combeite + Na_2Ca_2Si_2O_7ss$	Fig. 5E
15	$Liquid + Na_2CaSiO_4ss + Na_2Ca_6Si_4O_{15}$	Fig. 5F
16	$Liquid + Na_2Ca_2Si_2O_7ss + Na_4CaSi_3O_9ss$	Fig. 5G
17	$Liquid + Na_2Ca_2Si_2O_7ss + Na_2CaSiO_4ss$	Fig. 5H
18	$Liquid + CaSiO_3 + Ca_3Si_2O_7$	Fig. 5I







Fig. 14. Computed x, T diagram of the Ca_2SiO_4 -Na₄SiO₄ section by FToxid database [20].



Fig. 15. Computed isothermal sections of the Na_2O -CaO-SiO₂ system at 1000 °C by FToxid and MTOX databases [20,24].



Fig. 16. Computed isothermal sections of the Na_2O -CaO-SiO₂ system at 1200 °C by FToxid and MTOX databases [20,24].

3.7. Primary phase field of combeite

As can be seen in Fig. 13, the combeite phase obtained was located in the pseudo-binary section of Na₂SiO₃ and CaSiO₃, ranging for CaSiO₃ concentration from 50 to 73 mol-%. The solubility of CaSiO₃ in combeite increased with increasing temperature when it in equilibrium with CaSiO₃. This observation agrees with the computational phase equilibria of the FToxid database [20] and the data by Moir and Glasser [14]. The liquidus point data obtained in the present investigation are in good agreement with the FToxid database [20] between 1000 and 1200 °C and the MTOX database [24] at 1000 °C (Figs. 15 and 16). MTOX database [24] regards combeite as a stoichiometric compound of Na₂Ca₂Si₃O₉. Its computed liquidus line at 1200 °C (Fig. 16) is in poor



Fig. 17. Computed isothermal sections of the Na₂O-CaO-SiO₂ system at 1400 °C by FToxid and MTOX databases [20,24].



Fig. 18. Evaporation behavior of Na_2O in the Na_2O-CaO-SiO_2 system at 1200 $^\circ\text{C}.$

agreement with the data obtained in the present investigation. Liquid compositions with the double saturation of combeite with Na₂Ca₃Si₆O₁₆ (Fig. 5B), Na₄CaSi₃O₉ss (Fig. 5C), CaSiO₃ (Fig. 5D), and Na₂Ca₂Si₂O₇ss (Fig. 5E) were obtained in the present investigation. It means that the locations of the intersections and phase boundaries between combeite with the mentioned primary phase fields are now known in the diagram for the temperature range between 1000 and 1200 °C.

3.8. Primary phase field of Na₂Ca₂Si₂O₇ss

The present study indicates that Na₂Ca₂Si₂O₇ is not a stoichiometric compound but forms a solid solution, Na₂Ca₂Si₂O₇ss, for which the Na₂O concentration in the solution varies from 17.6 to 26.5 mol-% at 1200 °C. This finding provides novel experimental verification showing that Na₂Ca₂Si₂O₇ss can form a solid solution. MTOX database [24] considered Na₂Ca₂Si₂O₇ as a stoichiometric compound, and the computed liquidus line at 1200 °C is in poor agreement with the data obtained in the present investigation (Fig. 16). FToxid database [20] predicted that Na₂Ca₂Si₂O₇ss primary phase field does not exist at 1200

Table 6

Invariant points and the respective phases obtained in present study, pervious investigation [19] and computation [20] by FToxid database.

Point	Phases	Туре	Temperature,	Liquid (mol-%	Composit	ion,
			°C	Na ₂ O	CaO	SiO_2
A	$L + Ca_2SiO_4 = Ca_3Si_2O_7 + Na_2Ca_2Si_4O_2 = Ca_3Si_4O_2 = Ca_4Si_4O_2 = Ca_4Si_4$	Peritectic	1295	7.2	49	43.8
В	$\begin{split} L &= Ca_3Si_2O_7 + \\ Na_2Ca_6Si_4O_{15} + \\ Combeite \end{split}$	Eutectic	1285	13.3	41	45.7
	$L = CaSiO_3 +$		1280			
С	Combeite + Ca ₃ Si ₂ O ₇ L + Na ₂ CaSiO ₄ ss	Eutectic	1400 [19] 1287.7 [20]	11.3	39	49.7
D	$= Na_2Ca_2Si_2O_7ss$ + Na_2Ca_6Si_4O_{15}	Peritectic	1275	29.7	28.1	42.2
F	$Na_2Ca_6Si_4O_{15} =$ $Na_2Ca_2Si_2O_7ss +$ Combeite	Peritectic	1220	20.3	34.4	45.7
G	$\begin{array}{l} L+Combeite=\\ Na_2Ca_2Si_2O_7ss+\\ Na_4CaSi_3O_9ss \mbox{ or }\\ L+\\ Na_2Ca_2Si_2O_7ss=\\ Na_4CaSi_3O_9ss+\\ Combeite \end{array}$	Peritectic	1130	34.4	20.3	45.3
	L + Combeite =		1050			
Н	$Na_2Ca_3Si_6O_{16} + CaSiO_3 \text{ or}$ $L + CaSiO_3 = Na_2Ca_3Si_6O_{16} + CaSiO_16 + CaSiO_2 = Castoria + Ca$	Peritectic	1046.1 [20]	18.8	15.6	66.6
I	$L + CaSiO_3 = Na_2Ca_3Si_6O_{16} + SiO_2$	Peritectic	1030 1027.9 [20]	13.3	12.5	74.2



Fig. 19. A schematic diagram showing the steps in the manufacturing of the glass-ceramics [28].

 $^{\circ}$ C (Fig. 16). Liquid at double saturation of Na₂Ca₂Si₂O₇ss with Na₂Ca-SiO₄ss (Fig. 5H) was observed in the present investigation and, thus, the location where these two primary phase fields cross each other can be located in the ternary diagram.

3.9. Primary phase field of Na₂CaSiO₄ss

The details of the location of Na₂CaSiO₄ss in the diagram are presented in Fig. 14. As can be seen in Fig. 14, Na₂CaSiO₄ss is stable in the pseudo-binary Na₄SiO₄ and Ca₂SiO₄ systems. In the ternary diagram, the solid solution is located at a constant SiO₂ concentration of 33.33 mol-%. Between 1100 and 1400 °C, the maximum solubility of Na₄SiO₄ concentration in the solid solution is between 55.8 and 66.4 mol- % and it



Fig. 20. Liquidus contours and primary phase fields of Na₂O-CaO-SiO₂ system.

increases with increasing temperature. FToxid [20] and MTOX databases [24] regard Na₂CaSiO₄ss as a stoichiometric compound in their assessments. Thus, the computed liquidus at Na₂CaSiO₄ saturation is in poor agreement with the data obtained in the present investigation. As can be seen in Fig. 15, at 1200 °C, the FToxid database [20] predicts that the ternary liquid can contain CaO up to around 53 mol-%. In contrast, according to the present investigation, the maximum CaO concentration in the ternary liquid is only 27.2 mol-%. This can be explained by Na₂CaSiO₄ss being stable within a wide composition range. Thermodynamically, it will reduce the fully liquid region in the ternary diagram. Therefore, the insertion of Na₂CaSiO₄ss to the thermodynamic evaluation of the Na₂O-CaO-SiO₂ system will significantly change the phase assembly of the computed phase diagrams. Accordingly, it is highly recommended that this phase is taken into account as the diagrams in this ternary system are employed in many industrial applications.

3.10. Primary phase field of Na₄CaSi₃O₉ss

It can be seen in Fig. 15 that Na₄CaSi₃O₉ss is stable at 1000 °C and the liquidus contours obtained in the present investigation are in good agreement with FToxid [20] and MTOX database [24] calculations. Na₄CaSi₃O₉ss (Fig. 13) was also observed to form a solid solution in a narrow range of compositions, as also reported by Moir and Glasser [12]. Since the liquidus contour at double saturation of Na₄CaSi₃O₉ss and Na₂Ca₂Si₂O₇ss (Fig. 5G) was analysed in the present investigation, these solid solutions will be in equilibrium with each other along the univariant lines at different temperatures. However, the FToxid database [20] predicts that the Na₄CaSi₃O₉ss primary phase field does not connect to the Na₂Ca₂Si₂O₇ss field, but to Na₈Ca₃Si₅O₁₇, as reported previously by Segnit [6].

3.11. Primary phase field of Na₂Ca₃Si₆O₁₆

It can be seen in Fig. 15 that the liquidus data at $Na_2Ca_3Si_6O_{16}$ saturation obtained at 1000 °C are in good agreement with the computation using the FToxid database [20]. In contrast, the MTOX database [24] predicts the liquid phase domain at $Na_2Ca_3Si_6O_{16}$ saturation to be narrower than the data obtained in the present investigation (Fig. 15).

3.12. Invariant points

By definition, a invariant reaction has zero degrees of freedom in given conditions. Thus, it is almost impossible to determine the exact location of an invariant point at a certain total pressure and temperature in the ternary diagram by experimentation, as the starting mixture already fixes the compositions of the equilibrium phases. Thus, the locations of invariant points were interpolated from temperature and composition data in the ternary diagram. The solid phases analyzed are plotted in Fig. 12 to examine the Alkemade lines [26,27] used to determine the type of the invariant point. The Alkemade lines connecting two crystalline compounds were drawn as solid red lines. The lines can only be drawn if the phase fields of the two crystals intersect each other, i.e., they share a common univariant line. Fig. 12 shows the lines of intersection between two adjacent primary phase fields, drawn as black solid lines with arrows indicating the direction of decreasing temperature.

Fig. 12 shows the Alkemade lines of $Na_2Ca_6Si_4O_{15}$ - Ca_2SiO_4 , $Na_2Ca_6Si_4O_{15}$ - Na_2CaSiO_4 , $Na_2Ca_6Si_4O_{15}$ - $Na_2Ca_2Si_2O_7$, $Na_2Ca_6Si_4O_{15}$ - $Na_2Ca_2Si_2O_7$, $Na_2Ca_6Si_4O_{15}$ - $Na_2Ca_2Si_2O_7$, $Na_2Ca_6Si_4O_{15}$ - $CaSiO_3$, SiO_2 - $Na_2Ca_3Si_6O_{16}$, combeite- $Na_2Ca_3Si_6O_{16}$, combeite- $CaSiO_3$, combeite- $Na_2Ca_2Si_2O_7$, combeite - $Na_2Ca_6Si_4O_{15}$, combeite- $Na_4CaSi_3O_9$ ss, combeite - $Ca_3Si_2O_7$, $Na_2Ca_6Si_4O_{15}$, $CaSiO_3$, and $Ca_3Si_2O_7$ - Ca_2SiO_4 . $Na_2Ca_2Si_2O_7$, Ca_3SiO_2 - $Na_2Ca_2Si_2O_7$, Ca_3SiO_2 - $Na_2Ca_2SiO_7$, $CaSiO_3$, and $Ca_3Si_2O_7$ - Ca_2SiO_4 . $Na_2Ca_2Si_3O_9$ is used to describe combeite because the solid solution composition of combeite will approach to the fixed composition, $Na_2Ca_2Si_3O_9$, upon cooling. This similar approach of composition change of the solid solutions was also applied for Na_2CaSiO_4 ss, $Na_2Ca_2Si_2O_7$ ss, and $Na_4CaSi_3O_9$ ss.

Three Alkemade lines inside the ternary system cross their respective boundary lines of two primary phase fields. Firstly, the boundary between primary phase fields of combeite and Ca₃Si₂O₇ crosses the Alkemade line of combeite and Ca₃Si₂O₇ at point X₁. Secondly, the Alkemade line of combeite and Na₂Ca₆Si₄O₁₅ crosses the boundary between the fields of combeite and Na₂Ca₆Si₄O₁₅ at point X₂. Thirdly, the boundary between the primary phase fields of combeite and CaSiO₃ crosses the Alkemade line of combeite and CaSiO₃ at point X₃. Thus, according to the Alkemade theorem [26,27], points X_1 , X_1 and X_3 must be the local maximum points at which temperatures start to decrease in two opposite directions in the lines connecting two adjacent fields. It means that B is a new ternary eutectic of liquid + combeite + $Na_2Ca_6Si_4O_{15} + Ca_3Si_2O_7$ located at the temperature close to 1285 °C. Point C is the ternary $eutectic\ point\ of\ liquid + CaSiO_3 + combeite + Ca_3Si_2O_7\ located\ close\ to$ 1280 °C. FToxid database [20] calculated Point C to be 1287.69 °C, although it was predicted as peritectic, not as eutectic point. However, Z. Zhang et al. [19] estimated it to be close to 1400 °C.

Furthermore, the present study suggests the existence of the following peritectic points: (1) liquid + $Ca_3Si_2O_7$ + $Na_2Ca_6Si_4O_{15}$ + Ca_2SiO_4 (Point A) close to 1295 °C; (2) liquid + $Na_2Ca_2Si_2O_7ss$ + $Na_2Ca_6Si_4O_{15}$ + $Na_2Ca_6Si_4O_{15}$ + $Na_2Ca_6Si_4O_{15}$ + $Na_2Ca_6Si_4O_{15}$ + $Na_2Ca_6Si_4O_{15}$ + $Ca_2Si_2O_7ss$ + $Na_2Ca_2Si_2O_7ss$ + $Na_2Ca_6Si_4O_{15}$ + $Ca_2Si_2O_7ss$ + $Na_2Ca_6Si_4O_{15}$ + $Ca_2Si_2O_7ss$ + $Na_2Ca_2Si_2O_7ss$ + $Na_2Ca_2Si_2O_7ss$ + $Na_4Ca_2Si_3O_9ss$ + combeite (Point G), located 1130 °C.

Peritectic points of liquid + Na₂Ca₃Si₆O₁₆ + CaSiO₃ + combeite (Point H) at 1050 °C, and liquid + Na₂Ca₃Si₆O₁₆ + SiO₂ + CaSiO₃ (Point I) at 1030 °C are suggested in the present study. Points H and I were calculated using the FToxid database [20] to be at 1046.11 and 1027.88 °C, respectively. The invariant points and their phases are summarized in Table 6. Furthermore, Figs. 11 and 12 show that the ternary compounds of Na₂Ca₆Si₄O₁₅, Na₂Ca₂Si₂O₇ss, Na₄CaSi₃O₉ss and Na₂Ca₃Si₆O₁₆ are located not in their own phase field, meaning that they melt incongruently. As can be seen in Figs. 11 and 12, crystals of Na₂Ca₆Si₄O₁₅, Na₂Ca₂Si₂O₇ss and Na₂Ca₃Si₆O₁₆ are located in the fields of Ca₂SiO₄, Na₂Ca₆Si₄O₁₅, combeite and CaSiO₃, respectively.

3.13. Implications of the results on the glass and ceramic manufacture

A typical manufacturing process of glass-ceramics is presented schematically in Fig. 19 [28]. It involves melting an oxide mixture in a refractory-lined container to homogenize the melt. The viscous melt is formed to the desired shape, after which the sample is thermally treated to nucleate (T_N , Fig. 19) the desired crystalline phases. Then, the temperature is increased to value providing growth of the crystals (T_G , Fig. 19). Finally, the glass-ceramic sample is annealed to room

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temperature. The melting container size can be as large as an Olympic swimming pool if melting, for example, 1600 tons per day. Depending on the original mixture composition, the melting temperatures can be as high as 1600 °C [28] to produce a melt with a low enough viscosity needed to eliminate the gaseous inclusions and dissolve all the crystal-line species in the original mixture.

For a glass-ceramic in the ternary system Na₂O-CaO-SiO₂, the ternary invariant point and liquidus temperatures can be employed to estimate the commencement of melting (0% L) and the exact melting point of the mixture (100 % L). The data generated in the present work can also be used to figure out the crystalline phases that form in the thermal treatments of glass-ceramics. However, it should be pointed out that the actual melting temperatures of viscous melts for glasses and glassceramics are also affected by the fining and homogenizing criteria.

Fig. 20 shows the liquidus contour compared with the values based on the FTOxid [20] database. The compositions of typical glasses for daily applications e.g., for windows and container glass are also marked in the figure as a large grey circle [28]. Information about the eutectic points is critical for designing the glass since it quantifies the compositions which fully melt at the lowest temperature. This information might be useful when designing compositions with reduced energy and manufacturing costs. Fig. 20 shows that the melting point of a typical glass is actually lower if predicted by the data obtained in the present study than those predicted by FTOxid [20]. The difference is around 50 °C and increases up to 100 °C if the concentration of the mixture located in the Na₂Ca₃Si₆O₁₆ primary phase field moves toward primary phase field of CaSiO₃ as it showed that liquidus of 1300 °C from present study combined with liquidus of 1400 °C from FTOxid [20]. Present data also can be used to predict the crystal phases and their proposition during annealing more accurately. For example, if the annealing is undertaken in sub-liquidus at certain temperature and as in the field of CaSiO₃ the liquidus surfaces obtained in the present study is lower it means that the proposition of crystal to liquid is also smaller than that predicted by FTOxid [20]. Thus, data obtained in present study can be used for process optimization during melting and annealing.

4. Conclusion

Equilibration-quenching-EPMA/EDS method suggested solid solutions of Na₂CaSiO₄ and Na₂Ca₂Si₂O₇ in a wide range of compositions in the system Na2O-CaO-SiO2. The impact of the solid solutions on the ternary diagram was investigated together with the primary phase fields of SiO₂, Na₂Ca₃Si₆O₁₆, combeite, Na₄CaSi₃O₉ss, CaSiO₃, Na₂CaSiO₄ss, Na₂Ca₂Si₂O₇ss, Na₂Ca₆Si₄O₁₅, Ca₃Si₂O₇ and Ca₂SiO₄. Novel liquidus data at Na₂CaSiO₄ss, Na₂Ca₂Si₂O₇ss, and Na₂Ca₆Si₄O₁₅ saturation conditions are reported.

Five new invariant points were suggested based on the data obtained. The liquidus lines at saturation with two different crystalline phases as functions of temperature indicated the locations of the univariant lines as intersections between two adjacent primary phase fields. Comparisons of the data with previous investigations, assessment and computational results by FToxid database [20] databases and MTOX database [24] were also undertaken. Implications of current findings on the manufacturing of the glass and glass ceramics were discussed. The results suggest that Na₂O-CaO-SiO₂ system needs to be reassessed as the many new stable compounds are likely to significantly change many features of the phase properties and the computationally predicted phase equilibria. This study provides fundamental data to be included in the thermodynamic databases.

Data availability

Raw data and other supplementary material are available at the following repository: osf.io/7zpcj.

No data was used for the research described in the article. Data will be made available on request.

Declaration of Competing Interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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Appendix A

Isoplethal study

To evaluate the univariant line and its direction in the liquid projection diagram of Fig. A1, an isoplethal study was undertaken. In such a study, the reactions and phase relations taking place during cooling of a sample with constant total composition (isopleth) are examined [27]. Furthermore, the study can be used to evaluate how the composition of the solid solution changes during cooling. If the univariant line and the direction of decreasing temperature are correctly constructed, the phase relations obtained during cooling from the isoplethal study will agree with the experimental data.

Fig. A1 shows the line of V- L-P -R-N-G which is the path of the liquid composition changing when mixture V (19.9 mol-% Na₂O, 36.7 mol-% CaO, 43.5 mol-% SiO₂) is cooled from above liquidus to room



Fig. A1. Isoplethal study and cooling paths of mixtures V on the ternary diagram of Na₂O-CaO-SiO₂.



Fig. A2. Isothermal section at room temperature [23,24].

temperature at which all the reactions are completed. When temperature touches the liquidus temperature of 1345 °C, Na₂Ca₆Si₄O₁₅ is the first solid to form as V is located in the primary phase field of Na₂Ca₆Si₄O₁₅ and the liquidus composition will follow the cooling path of V-L during further cooling. When the cooling path touches the boundary line of D-F at point L, Na₂Ca₆Si₄O₁₅ starts to be reabsorbed and a crystal Na₂Ca₂Si₂O₇ss starts to form, and the cooling path now follows line of L-P to the direction of peritectic point F. From L to P, Na₂Ca₆Si₄O₁₅ is reabsorbed and Na₂Ca₂Si₂O₇ss is crystalized, simultaneously. At P, as the tie line Na₂Ca₂Si₂O₇ss the field of Na₂Ca₂Si₂O₇ss to point R in the boundary line of F-G without reaching point F. At temperature of P, Na₂Ca₂Si₂O₇ss start to form and just below P, Na₂Ca₆Si₄O₁₅ disappears. Cooling path now follows the line of R-N on which combeite



Fig. A3. Phase transformations of sample V during cooling from the melt to 1200 $^{\circ}$ C and further to room temperature, respectively.

and Na₂Ca₂Si₂O₇ss crystalize simultaneously from the liquid. When temperature of 1200 °C is reached, the composition of the liquid is represented by point N and the solid phases are combeite and Na₂Ca₂. Si₂O₇ss containing 21.5 and 17.3 mol-% Na₂O, respectively (the equilibrium composition of sample NCS-107). They are not stoichiometric compositions of Na₂Ca₂Si₃O₉ and Na₂Ca₂Si₂O₇.

As the initial mixture of V is located in the Alkemade triangle of Na₂Ca₂Si₂O₇-Na₄CaSi₃O₉-Na₂Ca₂Si₃O₉, according to Alkemade rule, the final assemblage of the solids when all reactions are completed, must be composed of these three stoichiometric crystals with the final composition of V. Thus, during further cooling from 1200 $^\circ$ C, the path will follow the line of N-G and all reactions with liquid will end at the peritectic point of G. At point G, Na₄CaSi₃O₉ss start to form, and at a temperature just below G, all liquid vanished. However, the reactions continue during cooling from G to room temperature. The reactions now are in the solid phase and they stop when the three solid solutions have been transformed into stoichiometric compounds. During cooling, from point G to room temperature, combeite composition will shrink to Na₄CaSi₃O₉. When it approaches Na₂Ca₂Si₃O₉, it will exsolve Na₂O and SiO₂ which are then dissolved by Na₄CaSi₃O₉ss and Na₂Ca₂Si₂O₇ss forming stoichiometric of Na₄CaSi₃O₉ and Na₂Ca₂Si₂O₇, respectively. Therefore, when the solidification is completed the final crystal composition is 65.4 mol-% Na2Ca2Si2O7, 6.1 mol-%, Na4CaSi3O9, 28.6 mol-% Na2Ca2Si3O9 (point V). This is also indicated by the calculation by MTOX database [24] according to which the Na2Ca2Si2O7-Na4CaSi3O9-Na2Ca2Si3O9 triangle exists at low temperatures (Fig. A2) and even already at 1000 °C, as seen in Fig. 15. The reactions and phase relations during cooling of sample Y and V are summarised in Fig. A3.

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