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INFLUENCE OF TEMPERATURE ON ULTRASOUND ABSORPTION AND STRUCTURAL PROPERTIES OF MELTS

Ultrasonic melt processing attracts considerable interest from both academic and industrial communities as a promising route improving melt quality. The significance of this problem is predetermined by the matter liquid state problem. In paper the ultrasound absorption and propagation speed v_s temperature are measured using multiple groups of samples, each group heated to a different temperature. This paper summaries the results on the evaluation of ultrasound absorption and the propagation speed, for calculating the ultrasound propagation speed in solutions, studies of the nucleation, growth and fragmentation of particles in liquid melts. It has been proven that melts with semiconductor properties are micro-inhomogeneous due to the existence of clusters in their atomic matrix. These results provide valuable new insights and knowledge that are essential for upscaling ultrasonic melt processing to industrial level.

Keywords: Ultrasound absorption; micro inhomogeneous melts; atomic mass; ultrasound propagation speed; coordination number

1. Introduction

The application of power ultrasound during liquid to solid transformation is believed to be an effective way to improve the solidification microstructures and mechanical properties [1]. In fact, the entire arsenal of modern experimental and theoretical physics is connected to the research of the physicochemical behavior of melts [2]. Acoustic methods are the most promising among experimental methods for research of the matter liquid state. They are simple, reliable, highly sensitive to changes of matter structure and the interatomic interaction.

The results of this research make it possible to predict the melts elastic properties of simple substances and extend it to complex substances.

At present, the electrophysical, thermophysical, thermodynamic and viscous properties of liquid semimetals and semiconductors based on the electronics industry have been widely studied.

However, the ongoing research in the field of studying these properties is not sufficient to solve the problem of the liquid semimetals and semiconductors structure. It is also impossible to obtain an unambiguous result by only structural research. In this aspect, it is known that «modern acoustic research methods are a powerful tool for obtaining information about the structure

of melts and semiconductors» [3]. Melts and semiconductors were not widely also studied, since the high-temperature acoustic experiments technology with aggressive melts of semimetals and semiconductors complicated the research process [1-5]. Our research includes: development liquid semimetals and semiconductors structure model; experimental and theoretical studies of the propagation speed and absorption coefficient of ultrasound temperature dependences in liquid semimetals and semiconductors; regularities generalization of liquid metals, semimetals and semiconductors structural properties.

A hypothesis about the micro-inhomogeneous structure of liquids arose in connection with Stuart's research using X-rays in the 1920. Micro-inhomogeneity extends to any melts from alkali metals to semiconductors. But the hypothesis remained a hypothesis, since there were no interpreted experimental data. The experiments were performed with a solidification apparatus incorporated with ultrasonic generator. The propagation of elastic waves is associated with the fundamental properties of material media, including the mass of particles, their space, the bonds between particles of matter. These indicators, namely, the inertial factor (mass of particles), the spatial factor (volume per particle), the stiffness factor between particles (compressibility), are sufficient for a general description of elastic waves absorption and propagation speed [6].

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2. Main results

As is known, the absorption coefficient is determined by the Stokes-Kirchhoff formula in the general case [7]:

$$\beta = \frac{2\pi^2 f^2}{\rho v_s^3} \frac{4}{3} \eta + \chi \left(\frac{1}{C_V} - \frac{1}{C_P} \right)$$

χ is the thermal conductivity coefficient, C_V , C_P is the heat capacity at constant volume, pressure.

The Stokes-Kirchhoff formula will take the form after the replacement $\gamma = \frac{C_P}{C_V}$:

$$\beta = \frac{2\pi^2 f^2}{\rho v_s^3} \left[\frac{4}{3} \eta + \frac{\chi}{C_P} (\gamma - 1) \right] \quad (1)$$

We will transform formula (1) to inertial, coupling and spatial factors based on the parameters $\eta = \rho v$, v is the kinematic viscosity, $\frac{1}{\rho v_s^2} = \alpha_S$ is the adiabatic compressibility, $C_P = \frac{dQ}{dT} \frac{1}{m}$ is the melt heat capacity:

$$\frac{\beta v_S}{f^2} = 2\pi^2 \alpha_S \left[\frac{4}{3} \rho v + \frac{\chi}{\left(\frac{dQ}{dT} \right)_P} m (\gamma - 1) \right] \quad (2)$$

Let $\rho = \frac{m}{V} = \frac{N_A M_A}{N_A V_A}$ and $m = N_A M_A$, N_A is the Avogadro number, M_A is the atomic mass, V_A is the atomic volume,

$$\sigma = 2\pi^2 \alpha_S \left[\frac{4}{3} \frac{v}{V_A} + \frac{\chi}{\left(\frac{dQ}{dT} \right)_P} N_A (\gamma - 1) \right]$$

Then we rewrite equality (2) in the form:

$$\frac{\beta v_S}{f^2} = \sigma M_A \quad (3)$$

The obtained Eq. (3) makes acoustic parameters monitoring in simple substances melts more accessible. Experimental measurements monitoring by value using reference data [8-12] is shown in Fig. 1. We have established the correlation dependence between measurement results $\frac{\beta v_S}{f^2}$ and the parameter values M_A , where σ is a constant value for each group. Note: the bonding factors intragroup similarity not only observed in condensed bodies, but also diatomic molecules it.

The highest value of the dissociation energy is observed for five electrons in the outer shell, the most rigid bonds exist in diatomic molecules [13]. Local bonds become stronger (partial atomic association) with an increase in the number of external electrons in melts due to the electron redistribution. There is no redistribution of electrons in diatomic molecules, since they bond

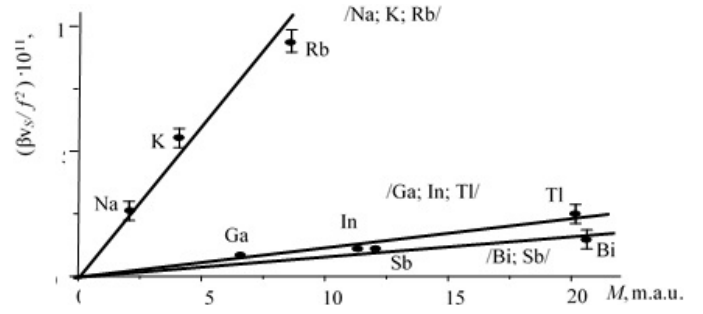


Fig. 1. $\frac{\beta v_S}{f^2} - M$ simple substances dependence at crystallization temperatures

with a single direction from one atom to another. If the number of external electrons is more than five in diatomic molecules, then individual electrons are transformed into a non-bonding state. This situation is explained by a decrease in the dissociation energy of diatomic molecules, starting from the oxygen group [13]. The closest packing coordination number (the number of bonds realization directions) can reach twelve in condensed matter. If there are more electrons than necessary to implement a uniform bond, they are redistributed. As a result, bonds more rigid are formed between individual neighboring atoms, associations of atoms are created.

It should be noted that the weakening of bonds between associates leads to an increase in bonds in associates and a decrease in the overall rigidity of the melt macroscopic volume. This is confirmed by the sulfur group example. There is a high correlation between the parameter $\frac{\beta v_S}{f^2}$ and the atom associates in the liquids of this group. The experimental values will be equal

$$\frac{\beta v_S}{f^2} = 19 \times 10^{-11} \text{ c for } S, \quad \frac{\beta v_S}{f^2} = 6,7 \times 10^{-11} \text{ c for } Se, \\ \frac{\beta v_S}{f^2} = 1,34 \times 10^{-11} \text{ c for } Te,$$

which indicates a decrease the atom associates in *S*, *Se*, *Te* melts.

We have shown in Fig. 1 that the parameter $\frac{\beta v_S}{f^2}$ increases from *Na* to *Rb* in the alkali metal series. This is confirmed by the fact that these groups of metals are prone to structure loosening with increasing atomic mass M_A . The $\frac{\beta}{f^2}$ temperature dependence is quite complex. Immediately after melting, there is a decrease in the absorption coefficient of ultrasonic waves, which increases monotonically with increasing temperature. This includes the alkali metals and further all other simple metals that are densely packed in the solid state.

Two-component liquid-metal solutions with monotonically increasing ultrasound absorption polytherms should also be attributed to this class. The absorption coefficient of ultrasonic waves decreases immediately after melting. In addition, the absorption coefficient of ultrasonic waves increases monotonically with increasing temperature. This includes the alkali metals

and further all other simple metals that are densely packed in the solid state. Melts, in which the absorption polytherms of ultrasonic waves do not increase monotonically with temperature, belong to the second class. This class includes semimetals and semiconductors, in which significant structural changes occur during melting. All melts with anomalous behavior of absorption polytherms and propagation rates (bismuth, antimony, tellurium, etc.) are characterized by the fact that their structural changes continue in a certain temperature range after melting.

This is especially pronounced in tellurium, antimony melts, etc. The polytherm $\frac{\beta}{f^2}$ anomalous behavior temperature interval of these melts coincides with the «after-melting» temperature interval. The experimental results on Eq. (3) are also can be used for compounds. For monitoring, let us consider GaSb and InSb compounds. Methods of descriptive statistics were used to systematize and describe the data. Experimental measurements monitoring by value using reference data [8-12] is shown in Fig. 1. The experimental results are shown in Figs. 2-12 for GaSb and InSb, Bi_{0,25}Sb_{0,75}, Bi_{0,5}Sb_{0,5}, Bi_{0,75}Sb_{0,25}, Bi-Sb, Sn_{0,30}Te_{0,70}, Sn_{0,5}Te_{0,5}, Sn_{0,70}Te_{0,30}, Sn-Te compounds. We used the TABLE 1 and TABLE 2 experimental measurements to build Figure with markers in Excel. The solid lines are the result of the experimental data approximation by the equation

$$v_s(T) = v_{SL} - \beta(T - T_L)$$

$\beta = \frac{dv_s}{dT}$ is the ultrasound propagation speed temperature coefficient, T_L is melting point, v_{SL} is ultrasound propagation speed at T_L .

Dependence $\beta = \frac{dv_s}{dT} - M$ is shown in Fig. 2 for these compounds.

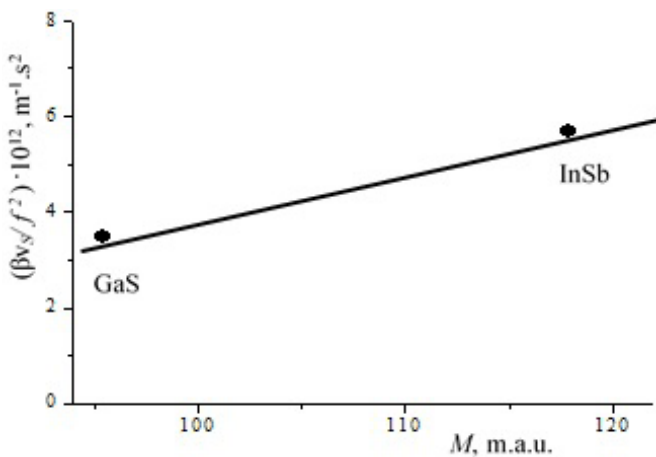


Fig. 2. Dependence $\frac{\beta v_s}{f^2} - M$ for GaSb and InSb systems

Thus, the acoustic analysis in electron melts experimental measurements shows that the absorption polytherms and the propagation velocity behavior of ultrasonic waves depends on the semimetals and semiconductors acoustic properties. This was the reason for separating these melts into an electron melts

separate class. A straight-line relationship between $\frac{\beta v_s}{f^2}$ and M_A inertial factor has been established.

For calculations, melt compounds were rotated by acoustically equivalent liquids from atoms of the same type by mass that satisfies the condition

$$M = M_1 X_1 + M_2 X_2 \tag{4}$$

M_1, M_2 – atoms mass; X_1, X_2 – components atomic fractions.

The experimental results on Eq. (3) are also can be used for compounds. Let us consider Bi-Sb compound. The experimental results are shown in Figs. 3-5 for Bi_{0,25}Sb_{0,75}, Bi_{0,5}Sb_{0,5}, Bi_{0,75}Sb_{0,25} compounds.

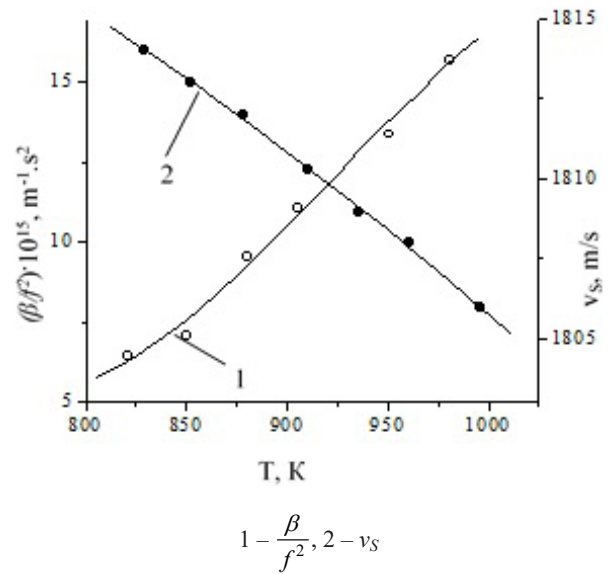


Fig. 3. Temperature dependence of ultrasonic absorption coefficient and propagation speed in the Bi-Sb system (composition Bi_{0,25}Sb_{0,75})

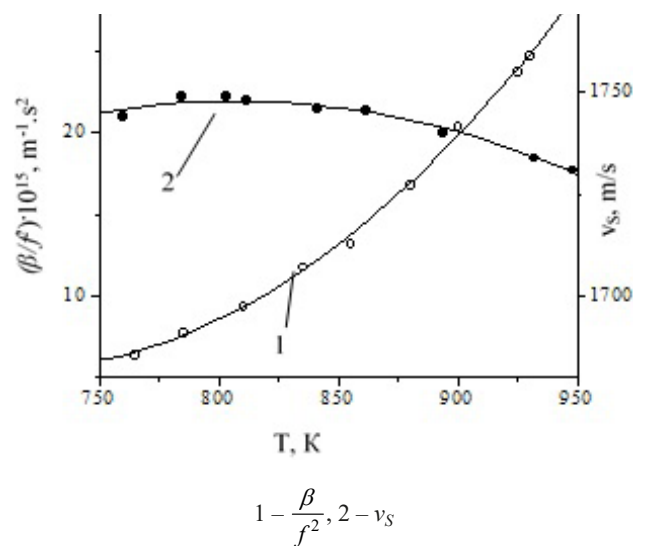
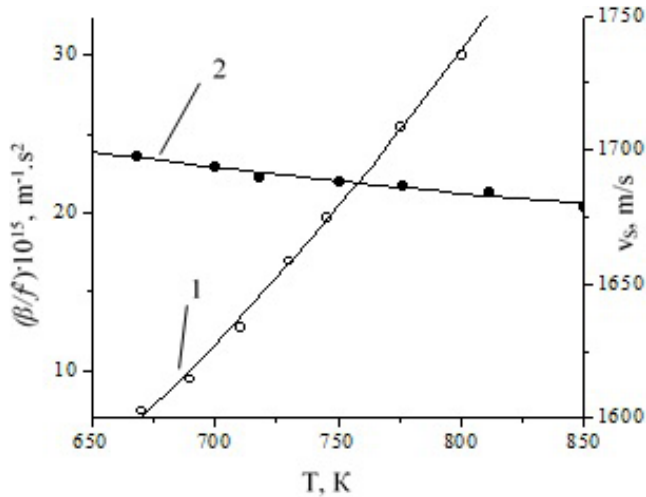


Fig. 4. Temperature dependence of ultrasonic absorption coefficient and propagation speed in the Bi-Sb system (composition Bi_{0,5}Sb_{0,5})

We have found that the ultrasound absorption coefficient increases with temperature in all Bi-Sb compositions. Dependence

Temperature dependence of Ultrasound absorption

Ga		Bi		Sb		Ge		Se		Te	
T, K	$\frac{\beta}{f^2} \times 10^{15}$	T, K	$\frac{\beta}{f^2} \times 10^{15}$	T, K	$\frac{\beta}{f^2} \times 10^{15}$	T, K	$\frac{\beta}{f^2} \times 10^{15}$	T, K	$\frac{\beta}{f^2} \times 10^{15}$	T, K	$\frac{\beta}{f^2} \times 10^{15}$
302	2,2	526	9,2	902	5,45	1211	1,56	520	725	734	14,5
325	2,9	545	9,3	918	5,28	1213	1,53	526	635	750	14,2
335	3,3	565	9,5	945	5,2	1220	1,5	544	630	762	13,8
370	3,5	599	9,8	960	5,44	1226	1,59	560	665	730	13,2
395	3,1	610	9,9	970	5,6	1229	1,6	574	565	812	13
405	3,4	615	10	986	5,8	1239	1,5	620	610	830	12,9
425	2,8	630	10,3	1000	5,83	1247	1,49	624	680	862	13
450	2	635	10,7	1020	6,4	1254	1,46	652	735	883	13,1
475	1,4	660	10,8	1035	6,5	1262	1,45	682	825	902	13,2
500	1,2	710	11,1	1050	7	1267	1,51	702	815	912	13,3
520	1,2	730	11,3	1078	7,3	1271	1,68	InTe		992	13,5
545	1,22	735	11,7	1094	7,5	1277	2,1	980	19,6	974	14,1
575	1,18	740	12	1104	7,8	1283	2,62	990	19,01	1006	15,1
585	1,28	755	12,4	1129	8,2	Ga ₂ Te ₃		948	18,5	1051	17
595	1,3	770	12,9	1138	8,9	1065	110,1	1010	18	1078	17,5
615	1,6	775	13,2	GaTe		1075	94,01	1025	17,5	1098	18,1
685	2,8	805	13,6	1100	62,1	1085	84,1	1040	16,9	1106	18,9
705	3,2	815	14,9	1112	56,2	1097	65,1	1050	16,6	In ₂ Te ₃	
745	3,6	820	15,4	1125	50,5	1110	55,2	1060	16,4	965	75,01
GaSb		835	15,9	1141	45,1	1112	50	1070	16,2	975	56,02
985	1,6	840	16,8	1167	38,1	1130	45,2	1085	15,8	905	42
1000	1,3	849	16,9	1178	35,8	1148	40,1	1097	15,5	1000	34
1020	1,2	855	17,3	1194	33,1	1160	38	Bi ₂ Te ₃		1015	26,2
1040	1,5	875	17,7	1204	28	1185	37	862	30	1030	22,1
1055	2,3	900	18,2	1228	24,2	1200	35,5	875	29,4	1040	21
1070	3	905	18,5	InSb		Sb ₂ Te ₃		890	29	1055	20,3
1077	3,8	910	19	814	2,78	902	24,4	900	28,5	1070	17
1090	4,4	915	19,5	820	2,65	909	23,1	930	27,4	1095	15,2
1105	5,4	935	20,8	840	2,9	923	22	956	26,5	Bi _{0,5} Sb _{0,5}	
1115	6,38	Sb ₂ Se ₃		855	3,1	933	21	978	25,8	765	6,4
Bi ₂ Se ₃		890	35,1	870	3,7	948	18,5	1009	25	785	7,8
1000	33	903	31,9	880	3,8	969	16,5	1042	24	810	9,4
1005	32,01	905	30	895	4,38	983	15	1080	23,4	835	11,8
1018	30,7	907	28	905	4,9	1007	14,1	Bi _{0,25} Sb _{0,75}		855	13,2
1025	30	912	27	920	5,4	1030	13,8	821	6,5	880	16,8
1037	29,3	915	25	930	5,98	1049	14	850	7,1	900	20,4
1045	28,8	928	22,6	935	6,2	1064	14,3	880	9,6	925	23,8
1049	28,7	935	20,8	950	6,6	1073	15	905	11,1	930	24,8
1060	28	940	20	969	7,3	Sn _{0,7} Te _{0,3}		950	13,4	In	
1070	27,8	955	21,5	SnTe		984	11,4	980	15,7	440	4,3
1074	27,7	965	19	1069	22,8	995	11,7	Sn		462	4,4
1090	27,5	975	18,9	1092	23,3	1020	12	500	4,8	500	4,8
1095	27,4	1000	18,1	1110	23,5	1045	12,2	560	5,4	520	5
Bi _{0,75} Sb _{0,25}		1019	18	1123	24	1050	12,7	584	5,1	570	5,9
670	7,5	Sn _{0,3} Te _{0,7}		1132	23,8	1080	14	619	5	598	6,9
690	9,5	868	18,5	1142	24,4	1087	14,5	680	6,5	625	7,3
710	12,8	890	18,4	1157	25,5	1111	16	740	9,5	640	7,8
730	17	910	19,5	1175	27,9	1116	17	820	10,5	675	8,6
745	19,8	938	20,6	1195	29			880	12	681	8,7
775	25,5	980	21					940	12,4	720	9,5
800	30,1	1020	23					980	12,6	740	10,2
670	7,5	1060	24,8					1000	13,4	770	11
		1080	26,8					1060	13,4		
								1120	15,2		



$$1 - \frac{\beta}{f^2}, 2 - v_s$$

Fig. 5. Temperature dependence of ultrasonic absorption coefficient and propagation speed in the Bi-Sb system (composition Bi_{0,75}Sb_{0,25})

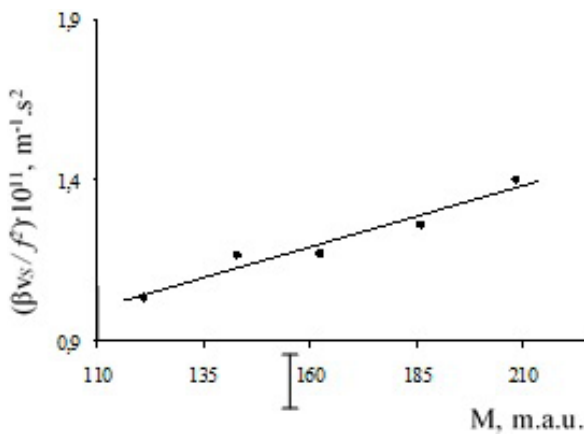


Fig. 6. Dependence $\frac{\beta v_s}{f^2} - M$ for Bi-Sb system

$\frac{\beta v_s}{f^2} - M_A$ is shown in Fig. 6 for these Bi_{0,25}Sb_{0,75}, Bi_{0,5}Sb_{0,5}, Bi_{0,75}Sb_{0,25} compounds. We approximated the experimental data with a linear function during their processing, the graph of which is a straight line. The linear function coefficients were calculated using the least squares method. The experimental data are practically placed on straight lines in Fig. 6. This confirms the closeness of the calculated indicators to the ideal ones. Indeed, the authors of [14] showed that the Bi-Sb system forms a regular solutions series with atoms random arrangement. The results in the coordinates $\frac{\beta v_s}{f^2} - M_A$ are shown for the Sn-Pb system in Fig. 7. We found that the $\frac{\beta v_s}{f^2}$ experimental measurements are practically placed on straight lines according to Pless's experiment. Note that the indicated results are typical only for ideal systems, that is, for melts with an atoms random distribution. Ideal solutions are solutions in which a negligibly small interaction between the particles of the constituent substances is assumed.

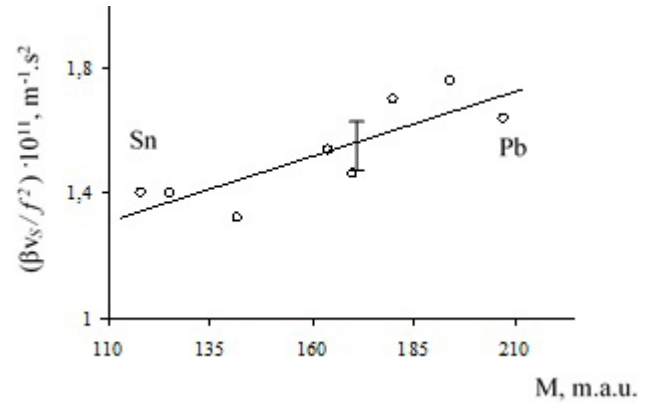
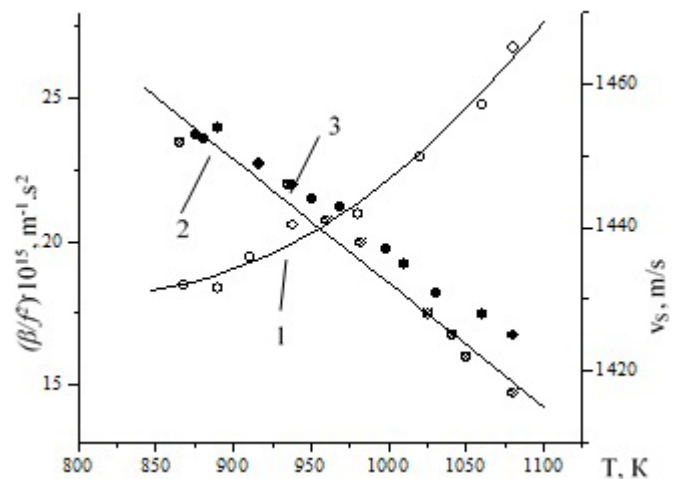


Fig. 7. Dependence $\frac{\beta v_s}{f^2} - M$ for Sn-Pb system [15]

We have developed an algorithm for determining the ideality or non-ideality degree of an arbitrary binary system based on monitoring the ultrasonic absorption coefficient and propagation speed experimental measurements. These are $v_s^2 - \frac{1}{M_A}$ and $\frac{\beta v_s}{f^2} - M_A$ diagrams.

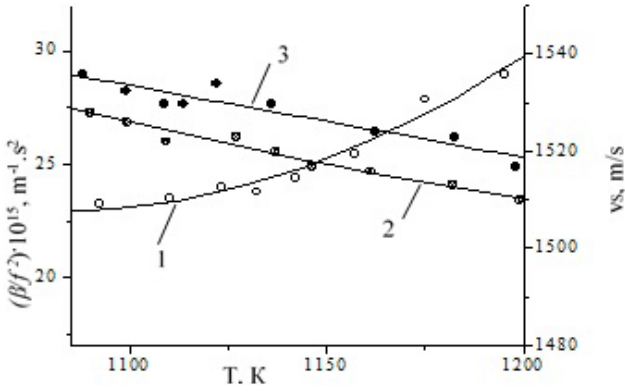
We study the Sn-Te system, which forms a congruently melting semiconductor chemical SnTe compound. The ultrasound speed and absorption experimental measurements in the compositions Sn_{0,30}Te_{0,70}, Sn_{0,5}Te_{0,5}, Sn_{0,70}Te_{0,30} are shown in Figs. 8-10. The ultrasound speed and absorption for liquid tin experimental measurements are shown graphically in Fig. 11.



$$1 - \frac{\beta}{f^2} \text{ the present work data, } 2 - v_s \text{ the present work data, } 3 - v_s [16] \text{ work data}$$

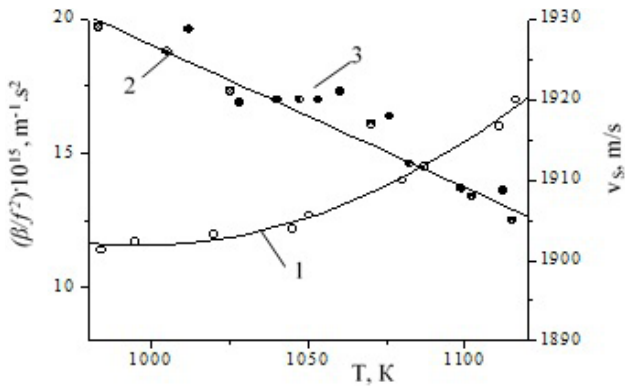
Fig. 8. Temperature dependence of ultrasonic absorption coefficient and propagation speed in Sn_{0,30}Te_{0,70} melts

We show a diagram for the Sn-Te system, it was previously studied by the [16,17] authors in Fig. 12. Calculations show that there are inconsistencies between the real isotherm and $\frac{\beta v_s}{f^2} - M_A$ linearity. Previously, for Bi-Sb and Pb-Sn systems,



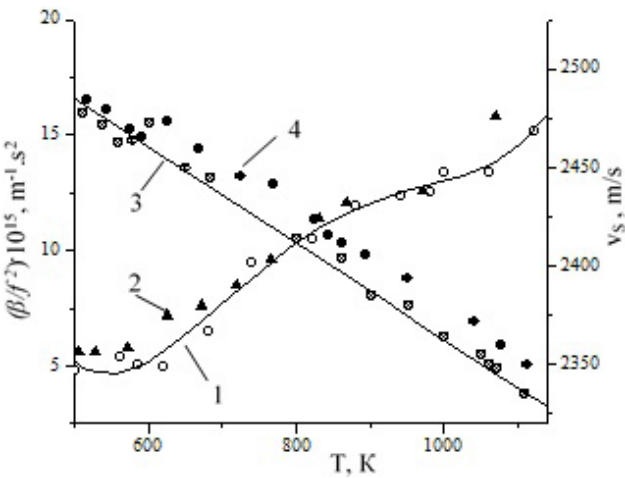
1 - $\frac{\beta}{f^2}$ the present work data, 2 - v_S the present work data,
3 - v_S [16] work data

Fig. 9. Temperature dependence of ultrasonic absorption coefficient and propagation speed in $Sn_{0.5}Te_{0.5}$ melts



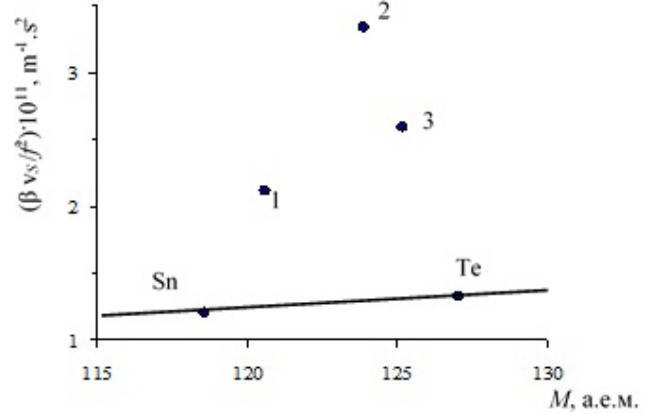
1 - $\frac{\beta}{f^2}$ the present work data, 2 - v_S the present work data,
3 - v_S [16] work data

Fig. 10. Temperature dependence of ultrasonic absorption coefficient and propagation speed in $Sn_{0.70}Te_{0.30}$ melts



1 - $\frac{\beta}{f^2}$, 3 - v_S the present work data, 2 - $\frac{\beta}{f^2}$ [16] work data,
4 - v_S [16] work data

Fig. 11. Temperature dependence of ultrasonic absorption coefficient and propagation speed in liquid tin



1 - $Sn_{0.30}Te_{0.70}$ 2 - $SnTe$ 3 - $Sn_{0.70}Te_{0.30}$

Fig. 12. Dependence $\frac{\beta v_S}{f^2} - M$ for Sn-Te system

it was found that these deviations are associated with melts microheterogeneity of intermediate concentrations. We have obtained a connection between the real isotherm $\frac{\beta v_S}{f^2} = f(M_A)$ and the corresponding linear diagram, which will allow us to evaluate the homogeneity or inhomogeneity of the melts and solutions. Consider a melt of the SnTe compound stoichiometric composition. We calculate the SnTe experimental measurement $\frac{\beta v_S}{f^2}$ and represent in Fig. 12 with the number 2. Through this point we draw a straight line parallel to the M axis. Let us assume that the SnTe melt is atomic and calculate the average M_A atomic mass by formula (4). An atomic SnTe melt corresponds to an extremely large M_A mass in an idealized system characterized by a linear dependence $\frac{\beta v_S}{f^2} - M_A$. As a result, the atomic mass of a real melt largely exceeds the average value atomic mass of the components. This unambiguously takes place in a situation where a real $SnTe$ melt consists not only atoms of tin and tellurium, but also larger particles, such as clusters or associates, whose masses exceed the component atoms masses.

To quantify statistical errors in the measurements we used simple descriptive statistics. The statistical error of a purely random sample is estimated by the formula:

$$\Delta = t \sqrt{\frac{\sigma^2}{n}}$$

Consider Ga as an example. The mean value is 2.272631579. The statistical or standard error is 0.215733269. The 95% confidence level is between 2 and 2.4. This means that if similar or random samples were re-selected, then about 95% of these samples would have average values between 2 and 2.4.

Ga		Ultrasound absorption	
Temperature		Mean	
Mean	502,7368421	Mean	2,272631579
standard error	30,41677766	standard error	0,215733269

Median	500	Median	2,2
Mode	#	Mode	2,8
Standard deviation	132,58366	Standard deviation	0,940359518
Sample variance	17578,4269	Sample variance	0,884276023
Kurtosis	-0,930109426	Kurtosis	-1,828842923
Skewness	0,217917553	Skewness	0,058423005
Rang	443	Rang	2,42
Minimum	302	Minimum	1,18
Maximum	745	Maximum	3,6
Sum	9552	Sum	43,18
Count	19	Count	19
Confidence level (95.0%)	63,90327858	Confidence level (95.0%)	0,45323878

We calculated statistical errors for all measurements from TABLE 3 and TABLE 4.

3. Conclusion

We used significance testing to compare results to determine if one is better than the other. Descriptive statistics were used using the parametric criterion (Student's t-test) to perform the analysis. Student's t-test can be used to determine if there are statistically significant differences between datasets.

Consider Ge as an example. Consider the results of the F-test to determine if there is a significant difference between the deviations. The results give three important values:

- F: ratio between dispersions.
- P ($F \leq f$) one-tail: the probability that variable 1 does not actually have a larger variance than variable 2. If it is greater than $\alpha = 0.05$, then there is no significant difference between the variances.

- F Critical one-tail: the value of F required for $P(F \leq f) = \alpha$. If this value is greater than F, it also means that there is no significant difference between the deviations.

F-test Two-sample for variance		
	Variable 1	Variable 2
Mean	1,660769231	1,649230769
Variance	0,111474359	0,111024359
Observations	13	13
df	12	12
F	1,004053165	
P($F \leq f$) one-tail	0,497262556	
F critical one-tail	2,686637112	

In our case $P = 0.497263 > \alpha = 0.05$, also $F_{cr.} = 2.686637 > F = 1.004053$. Therefore, there is no significant difference between the deviations.

We choose the appropriate T-test. As in the F-test, if $P(T \leq t)$ is greater than $\alpha = 0.05$, then there is no significant difference. However, in this case, two P values are given: one for the single-tailed test and the other for the double-tailed test. In this case, we use a double-tailed value, since any variable that has a larger mean will be a significant difference.

T-test: Two-sample assuming equal variances		
	Variable 1	Variable 2
Mean	1,660769231	1,649230769
Variance	0,111474359	0,111024359
Observations	13	13
Pooled variance	0,111249359	
Hypothesized mean	0	
df	24	

TABLE 3

Statistical errors of Temperature dependence of Ultrasound absorption

Compounds	Statistical errors	Compounds	Statistical errors	Compounds	Statistical errors
Ga	0,215733	Bi	0,670372	Sb	0,303635
Ge	0,092601	Se	27,1728	Te	0,482402
GaSb	0,582494	GaTe	4,298808	Ga ₂ Te ₃	7,758811
InTe	0,408283	In ₂ Te ₃	6,164237	Bi ₂ Se ₃	0,532322
Sb ₂ Se ₃	1,489689	InSb	0,440256	Sb ₂ Te ₃	1,146368
Bi ₂ Te ₃	0,735754	In ₂ Te ₃	6,164237	Bi _{0,75} Sb _{0,25}	2,991442
Sn _{0,3} Te _{0,7}	1,076162	SnTe	0,720425	Sn _{0,7} Te _{0,3}	0,666458
Sn	1,060004	In	0,62987		

TABLE 4

Statistical errors of Temperature dependence of Ultrasound propagation speed

Compounds	Statistical errors	Compounds	Statistical errors	Compounds	Statistical errors
Ga	4,873843	Bi	3,83035	Sb	1,242948
Sn	13,03025	Se	14,37275	Te	14,21959
In	9,527796	GaTe	2,080031	Ga ₂ Te ₃	3,708099
InTe	34,30029	In ₂ Te ₃	63,85878	Bi ₂ Se ₃	5,854248
Sb ₂ Se ₃	3,144983	SnTe	2,224209	Sb ₂ Te ₃	3,350349
Bi _{0,25} Sb _{0,75}	128,6236	Bi _{0,5} Sb _{0,5}	2,196026	Bi _{0,75} Sb _{0,25}	2,34702
Sn _{0,3} Te _{0,7}	4,236088	Sn _{0,7} Te _{0,3}	2,986577		

t-stat	0,088197409	
P(T <= t) one-tail	0,465225799	
t critical one-tail	1,71088208	
P(T <= t) two-tail	0,930451597	
t critical two-tail	2,063898562	

In our case $P = 0.930452 > \alpha = 0.05$, also $t_{cr.} = 2.063899 > t = 0.088197$. Therefore, our results are statistically significant.

This work demonstrated the feasibility and practical value of direct acoustic measurements in liquid metals. These measurements can be used for optimization of ultrasonic melt processing. The physical and chemical effects of ultrasound on liquid melts structure were investigated. In our study with specific components and ultrasound system, we determined that the chemical effect is an irreversible and permanent change in atom weight and the atom-weight distribution due to ultrasound. As the ultrasound intensity increases, the atom weight of liquid metals reduces and its atom-weight distribution becomes narrower; the orientation of liquid metals atom along the flow direction reduces (in melt state). Ultrasound vibration increases the motion of atom chains and makes them more disorder; it also affects the relaxation process of liquid metals, leading to weakening the elastic effect.

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