APPLICATIONS OF THE OPTICAL-DIFFERENTIAL IYERMAL ANALYGIS

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ABSTRACT

OOTA detects temperature contact-less, hence, it expands the upper usable temperature limit of DTA up to 3600°C accordingly suitable for studies of systems where enthalpic changes occur at high temperatures.

Application of the computer graphics for the evaluation of the OOTA curves is demonstrated on malting of Y₂O₃.Al₂O₃ and YAIO₃-GdAlO₃ **systems. Measurement errors occurring at temperatures above 1900°C are pointed out and possibilities for their eliminqtion or at least diminution are discussed.**

INTRODUCTION

The Optical Differential Thermal Analysis (ODTA)^{1,2}' detects **infrared radiation (IRl emitted by a sample heated inside the Black-Body** Cavity (BBC). The BBC fulfills a dual function, it is the temperature **source and the temperature reference stsndard.**

BBC is a perfect radiator which emissivity is unity. The sample emirsivity on the other hand, is a function of sample thermophysical and thermochemical properties, i.e., thermoradiative properties of the 9ample vary with the enthalpic change9 taking place In the sample. In the great majority of cases changes of enthalpy and emissivity are in concert thereby enhancing the resolution of the inset of ΔT_{max} **and/or** AT,*,. **peaks. This cooperation leado to a high 9ensitiUity of the ODTA which allows AT/t rates to be as low as 0.19Clmin. Consequently, phase relations can be studied at almost equilibrium conditions.**

ODTA APPARATUS

The OPTA apparatus consist9 of several parts: furnace with a power supply, furnace temperature control and a program system, two infra**r**
pyrometers and a data acquisition system. With the exception of the **furnace, most-of the ODTA components are commercially available** instruments. (The pyrometers are MAXLINE-SYSTEM M204, the furnace control and program system is MICRICON #82-300, manufactured by IRCON **Inc. and RESEARCH Inc., respectively. The signal acquisition system conristr of** IBM-PC **and IGY-AT.) For operation of the DDTA apparatus five Independent microprocessors are required. Each has different digital characteristics, therefore, signals. To elimin**e **digital signals are converted_to analog "cro9s talks", the O/A convertors ere mutually optically insulated. Numerour source9 of such instruments are available, hence, the selection of particular instrumentation depends on the operational range of the ODTA, and, on individual preference. The furnece is the critical part of the ODTA apparatus. The furnace design will be briefly discussed next.**

Thermal Ana&sis Proc. 9th ETA Congress. JerusaLm, Israel, 21-25 Aug. 1988 0040-6031188603.50 0 1988 Elscvier Science Publishers B.V.

The cross-section of the OOTA furnace, capable to operate in the **700 to 22OO*C range in a vacuum or an inert ger, i9 shown in fig.** 1.

Fig. 1. Left: Cross-section of the ODTA furnace: right: The graphite **Black-Body Cavity.**

A Black-Body Cavity b is derigned a% a hollow cylinder poritioned by the heat sink c_, concentrically with the heating element e. The heat aink is water-cooled as to provide sufficient heat removal from the BBC, especially at cooling rates higher than 1.0°C/min. The sample a is **placed inaide the BBC at ruch e height that it doea not interfere with the optical path of the radiometer d_,. The maximum wavelength varies over the OOTA temperature range. The wavelength of the IR radiation ir a nonlinear function of the temperature. Thus the IR radiation i9 converted to the temperature by a digital linearization using the Planck's law.**

Only few materials do not undergo enthalpic changes in the **1500-22OO*C temperature range. Among such materials are graphite and molybdenum, which both were found 9uitable for manufacturing of the BBC. A graphite BBC is ehown in Fig. 1 right. It In a hollow cylinder, which wall9 are corrugated to enhance the absorption of IR radiation. In the center of the bottom ir a sample holder, now replaced with the calibration reflector f. During mearurementr, the top of BBC 1% closed with a lid. The lid her a circular opening for mee9uring the sample** temperature with the pyrometer de. Pyrometer d, measures BBC temperature through an orifice in the side of the BBC (obscured in this view). **Diametera of both orifices are calculated not to interfere with the optical path of either radiometer.**

CALIBRATION

Two characteristics of optical pyrometers, the resolution power and

the ropestability, ~111 ultimately determine the accuracy of the OOTA. Optical pyrometers with the i2.C rerolutlon power and the ?l°C repeatability ere commercially aveileble. Yet, even such a good quality instrument will produce a low frequency noise in the OOTA. An attempt to filter much a noime by a common type of a filtar will necessarily dimlnlsh the accuracy and the sensitivity of 'the OOTA measurements. The meerurements are complicated by the imposaibillty to apply the Lambert's law acouretely, .and, to determine precisely the value of the shape factor A. Hence, these factors must be treeted reparately,end all such corrections included in a single correction factor.

Optical pyrometers are relative instruments and as such, they have to be calibrated. This is done by using materials with well known melting pointa. Two calibreted pyrometers are mounted on the OOTA furnace. Both are focused on the singular spot of reflector bloc f Fig. **1 right. The difference In temperature reedings between pyrometera d, and &, can be explained by the Lambert law. The difference is eliminated by setting the temperature reading of pyrometer de equal to** that of pyrometer d, stepwise over the entire temperature range.

A new tempereture difference will occur when the calibration reflector f is replaced by the sample holder. This time, the difference in the pyrometer readings is caused by a change of the shape factor A. **The temperature difference caused by the shape factor and by other Incalculable errors is called the integrated furnace background. It's value ir determined experimentally end used for calculation of** differential (DT) curves from the (T) curves.

INTEGRATED FURNACE BACKGROUND.

The heat transfer at temperatures over lOOO*C occurs predominantly by radfation. If both, the heater and the rample are at thermal equilibrium, there ir no heat trensfer either way. During thermal analysis, the sample temperature has to be Increased or decreased. In both the cases, a thermal gradieht between the heater and the sample must be eateblished. The degree of the gradient is determined by the AT/t rate and the shape factor A_. The degree of the gradient varies with temperature mainly due to the difference between the BBC and the sample emiseivity. It is also effected by the furnace atmosphere. Values of these factors are particular to the furnace, the GGC and the sample materials I.e. to the setup of the entire OOTA apparatus. How these factors affect a COT) curve la ahown on the [OT) curve of melting of YAlO= in Fig. 2.

For this reason, all the effectors are summarily determined and handled as an integrated furnace background (IFB). Due to the variability of these factors, the IFB has to be determined simultaneously with each individual measurement. For example, the (DT) curve shown in Fig. 2., now modified by the differentiation of the (T) curve with respect to the IFB, is shown in Fig.3.

SENSITIVITY AND REPRODUCIBILITY.

Yttrium and gadolinium aluminates form solid solutions over a large compositional range. Melting points of individual single phases differ only slightly. This makes it difficult to determine a phase diagram by the conventional methods. Dependence of the melting points of individual members on composition, i.e., a) YA10s; b) 90M% YAI0s 10M% GdA10s and c) 75M% YA103 25M% GdA103, is shown in Fig. 4.

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The lower end of the ODTA measurement range was calibrated on the melting point of copper, while the upper end on the melting point of a single crystel of sapphire. The accurecy of the ODTA measurement was found to be 12°C up to about 1700°C. Above this temperature, the error is larger. This is attributed to a particular malting behavior of sapphire'. The reproducibility of the measurements as determined from several hundred experiments is fi. end is depicted in Fig. S.

Curve a) in Fig. 5 is the same as in Fig. 3. Curve b) is from a sintered mixture of the powders. The apparent absence of an indication of an incongruent melting on curve b) is not attributed to the method, rather it shows that pre-melted samples give more detailed information.

CONCLUSIONS

Previously, Rupert^{3,} used optical measurements to determine (DT) and (dT) curves, but his approach was different from the ODTA, as he used non-black body conditions and extremely high AT/t rates. In any case, tha choice in methods is commanded by the specificity of needs.

Materials fmportant for ceramists and crystal growers are studied at temperatures not reachable by conventional DTA instruments. Therefore, at the present time, there is a lack of data for comparison with the data obtained by the DDTA. Distinct advantage of the DDTA is in the fact that it is not necessary to amplify the sensors' signals, i.e.
, the signals are not treated electronically. The only refinement of the ODTA data is mathematical. This approach allows to analyze the results using various aspacts of mathematical evaluation and, in case of uncertainty, it is again simple mathematically to modify the process.

The ODTA zero line or background is effected by numerous factors. Nevertheless, certain characteristics of the zero line contain a significant diagnostic information; hence, before any evaluation of the data is done, a careful study of the background data has to be made. Because the ODTA is a new technique, used in only one laboratory so far, there are no text books on its use and just few publications with which to confront the results. However, we believe that the ODTA is quite a potent technique, opening new approach for studying materials at high temperatures.

APPENDIX

An attempt is made here to refine the ODTA curves by a digital **filter in order to view the AT peaka in a grater detail. The digital** filter does not cause any shift along the abscissa. This approach is **aimed to the ODTA application to discern invariant and univariant points, i.e., to distinct a solid solution from an incongruently melting compound. At this time, however, there Is an insufficient experimental support for a positive proof.**

ACKNOWLGOGWENT

The author wishes to express his thanks to R. Allen and G. Bryant of the **YTL for their technical assistance and to E. Ramrden of the Boston University for the computer programmlng.**

REFERENCES

1 J. L. Carlavsky, Proceedinga of the 9th International Congress for Thermal Analyrir, Jerusalem, Israel, 1988.

- 2 J. L. Caslavsky, Report # MTL TR 88-11 U.S. Army Materials
- **Technology Laboratory Watertown, Yaarachusetts 02172-0001**
- **3 0. N. Rupert, Rev.: Scl. Inatr. 3%,1629 (1965).**

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