

Originally published as:

Müller, H.-J., Schilling, F. R., Lathe, C., Lauterjung, J. (2005): Calibration based on a primary pressure scale in a multi-anvil device - In: Chen, J. (Eds.), Advances in High-Pressure Technology for Geophysical Applications, Elsevier, 427-449

Title:	A standard-free pressure calibration in a multi-anvil device				
Authors:	Hans J. Mueller, Frank R. Schilling, Christian Lathe, Joern Lauterjung				
Plan:	Abstract Keywords1.Introduction2.Techniques, Methods and Materials Description2.1.Multi-Anvil High Pressure Apparatus MAX802.2.Ultrasonic Interferometry2.3.Determination of Sample Length2.3.1XRD-Scanning2.3.2.X-Radiography2.4.Experimental Procedures2.5.Gasket Insets - Anvil Support2.6.Samples3.Results and Discussion4.Conclusions Acknowledgments References TablesFigures and Figure Captions				
Corresponding Au	ithor:				
Mailing Address: Hans J. Mueller GeoForschungsZentrum Potsdam Department 4 Telegraphenberg Potsdam D-14473 Germany					
e-mail:	hans-joachim.mueller@gfz-potsdam.de				
phone:	+49/ 331/ 288-1443				
fax:	+49/ 331/ 288-1402				
Technical Base: Hardware: Operating System Text-Processor:	IBM compatible, Pentium-Processor Windows XP Professional WordPerfect 11.0				

A standard-free pressure calibration in a multi-anvil device

Hans J. Mueller, Frank R. Schilling, Christian Lathe, and Joern Lauterjung

GeoForschungsZentrum Potsdam, Dept. 4, Telegraphenberg, D-14473 Potsdam, Germany corresponding author's address: hans-joachim.mueller@gfz-potsdam.de

Abstract: A key question to all high pressure research arises from the reliability of pressure standards. There is some indication and discussion of an uncertainty of 10-20% for higher pressures in all standards. Independent and simultaneous investigation of the dynamical sound velocity and static compressibility by in situ X-ray diffraction (XRD) on a sample reveal the possibility of a standard-free pressure calibration and, consequently, an absolute pressure measurement. Ultrasonic interferometry is used to measure velocities of elastic compressional and shear waves in the multi-anvil high pressure device MAX80 at HASYLAB Hamburg. MAX80 enables XRD, X-radiography, and ultrasonic experiments. Two of the six anvils were equipped with lithium niobate transducers of 33.3 MHz natural frequency. NaCl was used as pressure calibrant, using the EoS of Decker (1971), and sample for ultrasonic interferometry at the same time. From the ultrasonic wave velocity data, v_p and v_s , we calculated the compressibility of NaCl as a function of pressure independent from NaCl-pressure calibrant. To derive the ultrasonic wave velocities from the interferometric frequencies of constructive and destructive interference require precise in situ sample length measurements. For a NaClsample this is of particular importance, because the sample is the most ductile part of the whole set-up. We measured the sample length by XRD-scanning and by X-radiography. The compressibility results, derived from the ultrasonic data, were compared with data of static compression experiments up to 5 GPa (Bridgman, 1940) and up to 30 GPa (Birch,

1986) using experimental data from Boehler & Kennedy (1980) and Fritz *et al.* (1971). At 1.2 GPa and 5.3 GPa the results of static compression data agree with our velocityderived compressibility data. In the range between 2 and 4 GPa our dynamical data have 1.5 - 3 % higher values. In general the pressure revealed according to Decker (1971) is in accordance to our standard-free pressure calibration. Consequently, up to 8 GPa the NaCl pressure standard has a reliability of at least 1%. However, at higher pressures the inaccuracy of the NaCl standard seems to exceed 1 %.

Key-words: pressure calibration, ultrasonic interferometry, compressional wave velocity, shear wave velocity, sodium chloride, multi-anvil high-pressure cell, X-ray diffraction, X-radiography.

INTRODUCTION

Multi-anvil devices are a very successful tool for experimental simulation of mantle conditions with relatively large samples. Accurate pressure determinations are critical to most high pressure measurements. However, pressure calibration and the reliability of pressure standards are discussed controversially.

The formation and the development of gaskets between the anvils causes a deviation between load per anvil surface and pressure inside the set-up because of friction, material variation of the pressure transmitting medium, minor fit variation in the set-up, minor adjustment variation of the set-up and the anvils to each other, and different compressibility of the samples. Recent pressure determinations in a gas piston-cylinder apparatus successfully reduced the uncertainty to 0.2 %, which is as low as that of free piston gages at 2.5 GPa (Getting, 1998). Therefore, in situ pressure measurements and precise standards are very important for this type of experiment.

Different options for pressure calibration exist

- using the known pressure of mineral reactions due to phase transitions, e.g., by measuring the change of electrical conductivity or using petrological experiments to determine mineral reactions. Several discrete measurements result in a pressure calibration curve (Luth, 1993),
- spectroscopic observation of a pressure dependent absorption band or peak, e.g., ruby chip (Piermarini *et al.*, 1975; Mao *et al.*, 1986)

(standard method for diamond anvil cells, not suitable for multi-anvil cells),

- continuous determination of the pressure dependent unit cell size of a standard by Xray diffraction, using the pressure marker's equation of state (EoS) (Decker, 1971; Chen *et al.*, 200).

The most common material to calibrate for conditions simulating the upper mantle is NaCl, following the equation of state published by Decker (1971), recently revised by Brown (1999). The other high-pressure calibrants like ruby-chips are usually calibrated against NaCl at low pressure. Progress in indirect pressure scale measurements has led to precision which exceeds the accepted uncertainty of the practical pressure scale by a factor of as much as five. A new indirect pressure scale would become available from the over-determination of the equation of state of a reference material by simultaneous X-ray and ultrasonic measurements (Ruoff *et al.*, 1973; Getting, 1998; Zha *et al.*, 1998).

MAX80 is a single-stage multi-anvil apparatus (Yagi, 1988) equipped for ultrasonic interferometry (Mueller *et al.*, 2002, 2003) and permanently located at HASYLAB,

Hamburg for having access to synchrotron radiation for in situ XRD measurements. We present simultaneous XRD- and high-pressure ultrasonic interferometry measurements of compressional and shear wave velocities of polycrystalline NaCl to determine a standard free pressure scale and to test the existing EoS by Decker (1971 and Brown (199). In situ sample length measurement, necessary for high precision ultrasonic interferometry, were performed by scanning both sample interfaces to the adjacent buffer and reflector and evaluating the XRD-spectra, as well as by X-radiography, i.e. taking X-ray shadow graphs of the set-up, recently installed at MAX80.

TECHNIQUES, METHODS AND MATERIALS DESCRIPTION 2.2. Multi-anvil high-pressure apparatus MAX80

MAX80 (Figure 1) is a single-stage multi-anvil high-pressure apparatus with six tungsten carbide anvils to compress a cubic sample volume of max. 8 x 8 x 8 mm. The anvils are driven by a 2,500 N uniaxial hydraulic ram, the top and bottom anvil directly, the lateral anvils by two load frames and four reaction bolsters, see Figure 2. Three anvilsets with different truncations exist - 6 mm, 5 mm, 3.5 mm. The maximum attainable pressures using 3.5 mm tungsten carbide anvils reach 12 GPa at 2,000 K produced by an internal graphite heater. The 6 mm truncation limits the maximum pressure to approximately 7 GPa.

Diffraction patterns are recorded in an energy dispersive mode (XRD) using white X-rays from the storage ring DORIS III at HASYLAB. MAX80 is equipped with a germanium solid state detector analyzing the diffracted white beam at a fixed angle with a resolution of 135 eV for 6.3 keV and 450 eV for 122 keV. Using a double-crystal, fixed-offset monochromator with silicon (311) single crystals, calibrated in the wavelength range of 0.4 to 0.6 Å, and a 2048 x 2048 pixels CCD-camera angle-dispersive X-ray diffraction (not used in this study) is also available.

The pressure is measured by energy-dispersive XRD using the high-pressure equation of state for NaCl (Decker, 1971). The method uses the observation of the elementary lattice cell compression of cubic NaCl crystals to derive the pressure in situ. These data are implemented in an in-house PC-program to calculate the resulting pressure at normal or given temperature. For details see Zinn *et al.* (1997), Vaughan (1993), and Shimomura *et al.* (1985).

The high pressure cell consists of a cube made of epoxy resin mixed with amorphous boron with the weight ratio 1:4 for better compressive strength containing the ultrasonic configuration, the heater, the pressure standard and the thermocouple. Although the graphite heater was not necessary for the experiments presented here it was not removed from the set-up for 6 mm anvil truncation to use the standard ultrasonic configuration of MAX80 (Mueller *et al.*, 2002). Removing the heater would result in pressure data not representative for the standard configuration. On the other hand, the 5 mm cube set-up was especially designed without heater to keep the sample surface bigger for stronger ultrasonic reflections, and to enhance the signal-to-noise-ratio. All interfaces between the sample and the close-fitting buffer rods / reflector bars are polished for optimal ultrasonic coupling (Figure 3). Additional coupling media were not used. Copper rings contact the heater at the top and bottom anvils. The sample is surrounded by rings made from boron nitride or glass ceramics for electrical insulation and as a quasi-hydrostatic pressure transmitting medium. Further details of the apparatus

-6-

are described on page, this volume and also by Mueller et al. (2002).

2.2. Ultrasonic interferometry

Ultrasonic interferometry, using the interference between the incident and reflected waves inside the sample, was first described by McSkimin (1950). Piezoelectric transducers for the generation and detection of ultrasonic waves are cemented at the polished rear anvils's side outside the true pressure cell. One or two of the original MAX80 anvil spacers (see Figure 2) were replaced by redesigned parts for ultrasonic experiments. The new spacers have a cavity in their center to keep the ultrasonic transducer free of any stress. In principle two types of ultrasonic set-ups were used in the presented experiments.

Asymmetrical set-ups are characterized by the optimization of buffers and reflectors, i.e. the buffer is made of a material resulting in intermediate acoustic impedance contrasts at both interfaces (anvil - buffer, buffer - sample) and the reflector material is selected for maximum reflection at the rear side of the sample. At ambient pressure the reflection coefficient for the NaCl-Pt interface is 80%, the reflection coefficient for the Pt-TC interface only 20%. This means, that only a minor amount is reflected between anvil and platinum, but most energy is reflected between NaCl and Pt, resulting in an optimized amplitude in the interference pattern. For the massive NaCl samples used in this study, powders were pressed, cut and re-machined. Buffers made from iron, aluminum, and Al₂O₃ ceramics were used. Platinum was found to be the optimum reflector material. To measure at one the velocity of compressional and shear

waves simultaneously with asymmetrical set-ups, requires the assemblage of both P- and S-wave transducers at one anvil or the use of a two-mode transducer as published by Kung *et al.* (2000). To ensure the maximum ultrasonic energy emission of the optimum cut transducers we used separate transducers for generation and detection, arranged in a circle as close as possible to one another (Figure 4). The geometrical error introduced by the eccentricity of 6.5 degrees is less than 0.5 %.

The other option - **symmetrical set-up**, i.e., buffer and reflector are made from the same material - requires ultrasonic measurements from the top and bottom anvil. Only one transducer for each wave type is concentrically assembled at one anvil's rear side. The advantage of this set-up is the optimum interference between direct and reflected waves because the transducer receives the reflected and interfered waves without any angular loss. On the other hand the symmetrical set-up results in additional energy losses due to non-optimum impedance contrasts between sample and buffer / reflector. For symmetrical set-ups we used platinum at both sides, which is an optimum reflector, but a poor buffer resulting in additional reflection losses, especially at the platinum-NaCl interface. In case of measurements at elevated temperatures, not performed in this study, only one of the "ultrasonic" anvils can be grounded. Even by using a dc-power supply small fluctuations of the current result in interference with the ultrasonic signals.

For generation and detection of the ultrasonic waves we used lithium niobate transducers, cold covered, overtone polished with a natural frequency of 33.3 MHz and a diameter of 5 mm. They were cemented at the polished rear anvil side using epoxy resin diluted by acetone to reduce its viscosity for minimizing the thickness of the glue film. This is of fundamental importance for the interferometric method to ensure rigid coupling to the anvil, because it requires a broad band characteristics of the transducer as

-8-

a result of strong attenuation. The ultrasonic anvils are equipped with one P-wave or Swave transducer, or with two couples of P-wave and S-wave transducers respectively, depending on whether an asymmetrical set-up and two-transducer method, or a symmetrical set-up and single transducer method was used. First tests were performed with 3.5 mm truncation anvils made from cubic boron nitride.

Figure 5 shows the electronic equipment for ultrasonic interferometry at MAX80. A PC-program controls the frequency sweep of the rf-generator by a frequency step of 100 kHz. An arbitrary waveform generator cuts wavelets (Shen et al., 1998) or double wavelets (Li *et al.*, 1998) of a duration of 20 ns to 4 μ s from the continuous sinusoidal signal of the rf-generator. The ultrasonic generator delivers the master trigger pulse and amplifies the received signal. For single transducer configurations, i.e., the transducer acts sequentially as generator and receiver of ultrasonic waves, a directional bridge is used to prevent the strong excitation wavelet from hitting the sensitive input of the receiving amplifiers. A power amplifier and pre-amplifier are used for samples with high damping or strong reflection losses at the interfaces. The multi-channel oscilloscope displays and digitizes the interference signals, finally stored on the PC's hard drive. The evaluation using an in-house computer program includes the selection and copying of the critical signal ranges, i.e., the buffer and sample reflections, their subtraction to isolate the interference between the signals, digital filtering, displaying the resulting periodic energy levels (constructive and destructive interferences) as a function of frequency, and finally displaying the resulting travel time curves as a function of frequency as well (Mueller *et al.*, 2003). The determined two-way travel time or its multiple inside the sample is represented by the bold straight line between the curves of opposite curvature in Figure 6.

2.3. Determination of Sample Length

The result of ultrasonic interferometric measurements is an equidistant sequence of critical frequencies for constructive and destructive interference of the reflected waves from the plane-parallel surfaces of the sample rod. Unfortunately the interference pattern does not only depend on the material properties of the sample, but also on sample length. Due to the sharp interference pattern, the travel time is determined with high precision better than 0.4 % - and the accuracy of the velocity determination mainly depends on the precision of the length measurement (Li et al., 2001). In situ sample length measurement in multi-anvil devices is not trivial, as it cannot simply be derived from measurements of the advance of anvils. Therefore sample deformation models, derived from direct length measurements prior and after the experiment (Knoche et al., 1997, 1998) are common usage, or it is assumed that the sample deforms purely elastically. The so called Cook's method (Cook, 1957) calculates the in situ sample length from the compressibility derived from measured elastic wave velocities. Our measurements with different samples in a variety of configurations showed that this assumption is only valid, if the sample is the strongest part of the buffer - sample - reflector combination. Knoche et al. (1997, 1998) had a hot isostatically pressed forsterite sample between two platinum buffer rods. Consequently the condition mentioned can be a good approximation for high-strength samples, as it was also the case in our experiments with San Carlos olivine, anorthite, clinoenstatite, and quartz (see Mueller et al., page ..., this volume), but not if a ductile sample is taken into account.

2.3.1 XRD-Scanning

Contrary to experiments with high-strength samples, a NaCl-specimen between Al₂O₃ ceramics or iron and platinum buffers is the most ductile part and will accommodate large parts of the total deformation (Mueller et al., 2003). Because sodium chloride deforms as a combination of ductile and elastic behavior, simple deformation models are not useful and measurements under in situ conditions are necessary. An advantage of ultrasonic measurements at a radiation source is that the sample length can be determined independently from the ultrasonic experiments. The first option is to scan the buffer - sample - reflector combination stepwise, crossing both the interfaces and determine the sample length by evaluating the in situ XRD-spectra (Figure 7). The circles represent the X-ray beam radius of about 50 μ m. The XRD-spectra close to the interface are a superposition of two spectra, because the X-ray beam penetrates both materials, i.e. Pt and NaCl, to some degree. The whole press (including the multianvil device) can be lifted by stepper motors with an accuracy of 1 μ m. By calculating the interface from the last and first pure spectrum the sample length can be determined much more precisely than the beam diameter is that is an accuracy of 5 to 10 μ m. An advantage of this method is, it requires no additional equipment and results in sufficient accuracy. The drawback is, it is highly time-consuming, about 20 minutes using the lowest step rate.

2.3.2 X-Radiography

Li et al. (2001) used X-radiography to measure the sample length in multi-anvil devices under in situ conditions, after the method was established in the nineties by other authors to observe under high-pressure conditions falling spheres in melts for viscosity measurements. At the latest when we introduced an ultrasonic data transfer function technique at MAX80 (see page, this volume) the XRD-scanning method was no longer adequate as the only available length measurement technique. A digital ultrasonic sweep for v_p and v_s lasts about 90 minutes. Consequently a duration of about 20 minutes for a XRD-scan of both interfaces was acceptable. But if the recording of two data transfer functions representing the whole v_p - and v_s -data requires only some seconds, the length measurement becomes the limiting factor.

As the first step to establish a X-radiography system the fixed double-slits unit of MAX80 was exchanged by an adjustable slits system. We used a four-blade highprecision slits system of ADC (Figure 8) equipped with four independent stepper motors including all the control electronics onboard. The maximum slits opening is 1 inch. The motion repeatability is 1 µm with a motion resolution of 0.4 µm. The MS Windows compatible IMS terminal software allows to control the slits system simply by the PC, already installed for the ultrasonic measurements. Because the four blades can be moved independently from each other the slits system is able to define the X-ray beam position and size. Differently from the original state the X-ray beam position can be controlled by the slits and the positioning table of MAX80 now. For X-radiography the blades are opened so far, that the X-ray beam covers the whole sample length including a part of the adjacent buffer and reflector rods. Using tungsten carbide anvils absorbing the synchrotron radiation (intense X-rays) the maximum vertical opening of the beam is adapted to the maximum available gap between the lateral anvils, of about 1.5 mm at normal pressure and less than 0.5 mm at maximum conditions. To limit the scattered radiation inside the hutch the slits is only opened to the size necessary for the sample length measurement.

First of all the X-radiography system (Figure 9) consists of an 0.1 mm thick Ce:YAG-crystal (by courtesy of IKZ) of 15 mm diameter in an adjustable aluminum mounting. It partially converts the X-ray shadow graph after passing through the set-up by fluorescence to an optical image of about 540 nm wavelength (light green), which is redirected to a CCD-camera by an aluminum-coated mirror. A beam-stop behind the mirror absorbs the non-converted X-rays. The Ce:YAG-crystal should be made as thin as possible to limit the warming up by X-ray absorption and to keep the optical image as sharp as possible, because the fluorescence creates optical images at all atomic planes inside the crystal. Extensive use of aluminum for X-ray exposed components is recommended to limit the warming of the parts by absorption. The decoupling of the optical image from the X-ray shadow graph by the mirror is necessary to prevent the CCD-camera from direct X-ray flux. The whole system is covered by a 2.5 mm thick Pbcasing for shielding from scattered radiation inside the hutch (Figure 10).

For taking images optimum for the following evaluation each shadow graph was recorded with three different exposition times, differing from each other by the four-fold exposition. The automatic exposure control failed because of the high intensity contrasts of the images. The evaluation of the shadow graphs is performed by densitometry profiling, i.e. the image processing software analyzes the brightness of the image along a pre-defined line. Figure 11 shows the shadow graph and the related image processing result for a NaCl-sample at 5 GPa pressure in linear and logarithmic scale. At the optimum exposition time the low-dense NaCl is displayed as pure white. The sample length, i.e. the number of zero density pixels at the central part of the image, is 149 pixels. Because of the small, but existing divergence of the X-rays, the shadow graphs and the sample have not necessarily the same size. Therefore the shadow graphs are calibrated, before the high pressure run starts, because at this time the sample length is exactly known from the preparation. From this calibration we know that the 149 pixels, displayed at Figure 11 represent a sample length of 1.94 mm. That means the accuracy is 1 pixel, i.e. 0.013 mm.

What are the accuracy limits for X-radiography? The principle limit is the wavelength of light, i.e. about 0.5 μ m. But in reality it gets worse, because the aperture of the objective is less than 0.5. To keep the camera outside the intensive X-rays, the working distance must be about 40 mm, very large, i.e. very disadvantageous for a micro objective. That limits the practical accuracy to about 1 μ m, what is a half order of magnitude better than X-ray scanning at the minimum. First results with a conventional consumer 5 Megapixels color camera with a minimum working distance of 70 mm demonstrate the potential of the used set-up and confirm the results of XRD-scanning. Because the image processing only uses the density of the image, first the color image is converted to a grey scale one. Therefore in the next experiments a 6 Megapixels black and white CCD-camera will be used at a working distance of 40 mm to guarantee a 1 μ m resolution.

2.4. Experimental Procedures

Three experiments were included in the evaluation. Run 2.2 is one of the first

ultrasonic experiments at all performed simultaneously with synchrotron XRDmaintenance in MAX80. Due to the similarity between the set-up of run 3.10 and 2.2 the sample deformation measured by synchrotron radiation during run 3.10 was used for both experiments. Set-up 2.2 had a buffer made of glass ceramics; set-up 3.10 had an iron buffer. Both asymmetrical set-ups had platinum reflectors. To make the pressure per load and deformation results comparable to other experiments the set-ups had a stepped graphite heater which was not in use during these experiments.

Run 3.27 used 6 cubic boron nitride (cBN) anvils with 3.5 mm truncation to increase the maximum pressure. Because cBN is an electrical insulator the rear side of the top and bottom anvil got a gold-platinum electrode for the transducers by sputtering. The top anvil was equipped with pairs of p- and s-wave transducers. In addition to that, the bottom anvil was equipped with a single p-wave transducer to compare the results of both configurations (see Figure 4). Due to electrical contact failure at the bottom piston only the symmetrical set-up with two platinum buffers could be used. The much smaller anvil truncation require boron-epoxy cubes of 5.5 mm length. To enlarge the reflection surface, i.e., to have a higher sample diameter a special set-up was designed without heater and insulator tube (Figure 3). The experiment showed that the friction between anvil's surface and gaskets was much higher than using tungsten carbide anvils resulting in a maximum pressure of 7.71 GPa.

2.5. Gasket Insets - Anvil Support

In normal use MAX80 forms the gaskets between the anvils from the boron epoxy

cube's material during the runs. This allows a simple and rapid sample change, ensures low X-ray energy loss by any additional materials and results in a good "high pressure efficiency", i.e., for a given load the pressures are relatively high, because the small gaskets formed by the cubes reduce the additional surface and hence, the "unproductive" part of the load. On the other hand a better lateral anvil support by an additional gasket results in a more homogeneous stress distribution inside the anvils leading to a higher maximum force to the sample cube and consequently higher pressures inside the set-up. Prefabricated gasket insets, normal for all double-stage multi-anvil devices, are a way for lateral anvil support at the expense of a lower pressure efficiency. For first tests we used gasket strips made from Klinger SIL C-4400 (Figure 12), an industrial sealing material made from NBR tied p-aramide fibres for tungsten carbide and cubic boron nitride anvils. The post-experimental optical inspection of the tungsten carbide anvils showed that the material starts to flow at the corners of the front face without any failure of the anvil. Because the gap width between the anvils was larger at elevated pressures the X-ray intensity was higher and the adjustment of the ray was easier. In other words, due to the reduced pressure efficiency the maximum pressure could be enhanced and thw XRD measurements could be improved. For cubic boron nitride anvils the friction between the gasket material and the anvils was too high, resulting in a stick-slip behavior and material failure. Material and shape of the gasket insets will further be optimized for future experiments.

The first experiments with tungsten carbide anvils showed ≈ 25 % higher maximum pressures compared to the standard MAX80 configuration because of increased lateral anvil support, reduced number of blow-outs, higher X-ray intensity, and a reduced probability of thermocouple cut-off during the experiments.

2.6. Samples

Sodium chloride powder of 99.5 % purity (analytical grade by Merck) was used as starting material. The medium grain size was 50 μ m. The powder was pressed to a sample cylinder of 10 mm diameter and a length of 20 mm using a load of 6 tons resulting in an effective pressure of 0.25 to 0.3 GPa. The millimeter sized samples (diameter 2.4 mm and 1.6 mm length for 6 mm anvil truncation, and diameter 3.1 mm and 1.1 mm length for 3.5 mm anvil truncation) for the high pressure experiments were shaped with a high-precision (\pm 0.5 μ m) cylindrical grinding machine and polished at the plane-parallel faces of the sample rod.

3. RESULTS AND DISCUSSION

The digitized interferometric signals stored on a PC's hard drive were processed using an in-house program. The resulting sequence of maxima and minima represents the frequencies for constructive and destructive interference. Picking all available maxima and minima as a function of frequency v allows the determination of the travel time τ inside the sample as the regression result for the horizontal point sequence between the curves of opposite curvature (Figure 6). The curvature is the result of an inappropriate use of the order of interference *n* according to $\tau = n 1/v$.

The calculation of wave velocities requires the sample length as a function of pressure. Consequently the precise measurement of sample deformation during the

experiment is essential for the accuracy of the whole method, because for higher degrees of deformation this contribution to the critical frequency interval can be higher than that of the change of sample's elastic properties. Figure 13 is the plot of v_p and v_s for the three experimental runs. The results for our runs are in agreement with previous results published by Frankel *et al.* (1976) within the limit of experimental errors (~1.5 %).

The velocities of run 3.27 are located between the values of the other two experiments and are used as average value. This run reaches the highest pressure and was used for further modeling.

The measured elastic wave velocities v_p (compressional wave) and v_s (shear wave) were used to calculate the adiabatic bulk modulus K_s and the corresponding compressibility κ_s .

$$K_{s} = \rho \left(v_{p}^{2} - \frac{4}{3} v_{s}^{2} \right) \tag{1}$$

and

$$\kappa = \frac{1}{K_S} \tag{2}$$

This calculation requires the density ρ of the sample as a function of pressure which is directly obtained by XRD measurements. In situ sample length measurement is the important basis, but in addition to that the study of the lateral deformation is also necessary. That becomes the more important, the less hydrostatic the pressure and the more ductile the sample is. Therefore a user of any multi-anvil device have to take care of set-up deformation. Different methods exist to meed the demands. The general form of the equation of state (EoS) is:

$$P(V,T) = P_1(V) + P_{TH}(V,T)$$
(3)

where P_{I} refers to the isothermal equation of state, and P_{TH} refers to the thermal pressure. For small compressions, the isothermal bulk modulus K_{T} can be approximated by:

$$K_{T} = -V \left(\frac{\partial P}{\partial V}\right)_{T} = K_{0} + PK_{0}' + \frac{P^{2}K_{0}''}{2} + \dots$$
(4)

Here K_0 , K_0 , and K_0 , are zero-pressure values of K and its first and second pressure P derivatives, at constant temperature. The first two terms usually suffice to represent ultrasonic measurements, but K_0 , appears to be negative and of a magnitude such that a quadratic in P leads to K = 0 (Birch, 1978). Therefore, only K_0 and K_0 , are used. Using published data for K, K and density at normal pressure ρ_0 (Birch, 1978, 1986; Holland and Ahrens, 1998) the density at given pressure can be calculated:

$$\frac{V}{V_0} = 1 - \kappa P \tag{5}$$

and

$$\rho_P = \frac{V_0}{V} \rho_0 \tag{6}$$

Another widely used approach is measuring and deriving the deformation of the sample from the ultrasonic experiment itself, called Cook's method (Cook, 1957; Kung *et al.*, 2001a, b).

$$S = 1 + \frac{1 + \alpha \gamma T}{3h_0} \int_0^P \frac{dP}{\left(\frac{1}{t_p^2} - \frac{4}{3}\frac{1}{t_s^2}\right)}$$
(7)

$$h_0 = 4\rho_0 l_0^2 \tag{8}$$

with S - linear compression

- α linear thermal expansion coefficient
- γ thermodynamical Grüneisen parameter
- *T* absolute temperature
- **P** pressure
- ρ_{θ} density at zero-pressure
- l_{θ} sample length at zero-pressure
- t_p travel time of compressional waves along the sample
- t_s travel time of shear waves along the sample

But this is only valid for

$$\frac{\rho}{\rho_0} = \left(\frac{l_0}{l}\right)^3 = S^3 \tag{9}$$

which means the deformation is purely hydrostatic, i.e., uniform in all directions of space. However our post-experimental examination of the set-up showed that this boundary condition is not achieved for our set-up and non-encapsulated NaCl-samples, because the sample is the most ductile part of the set-up. As a consequence of the gasket formation there is a reel-shaped deformation of the sample, i.e., the length decreases, the diameter at half the sample length slightly decreases or keeps constant, but the diameter at the front faces increases. Some minor parts of the material can be even squeezed out there. Therefore we used a more generalized equation published by Frankel *et al.* (1976). For a material whose equation of state is unknown, Katz and Ahrens (1963) showed that an equation of state can be solved for by assuming that the geometry of the specimen changes under pressure such that

$$\rho = \rho_0 X^n \tag{10}$$

where

$$X = \frac{l_0}{l} \tag{11}$$

with X - geometry characteristics

The parameter n is any positive number and is assumed to be independent of pressure. The change in the specimen density and thickness can be determined from the data as follows:

$$X^{n-2} = 1 + \left(\frac{n-2}{n}\right) \frac{1}{4l_0^2 \rho_0} \int_0^P Y dP$$
(12)

For $n \neq 2$, and

$$X = \exp\left[\frac{1}{8l_0^2 \rho_0} \int_0^P Y dP\right]$$
(13)

For n = 2, where

$$Y = \frac{1+\Delta}{\Delta f_p^2 - \frac{4}{3}\Delta f_s^2}$$
(14)

$$\Delta f_p = \frac{v_p}{2l} \tag{15}$$

$$\Delta f_s = \frac{v_s}{2l} \tag{16}$$

$$\Delta = \frac{9\alpha^2 T K_s}{\rho C_p} \tag{17}$$

with Δf_p - frequency interval between two critical frequencies for compressional waves

 Δf_s - frequency interval between two critical frequencies for shear waves

 C_p - specific heat at constant pressure

If the forces acting upon a specimen are perfectly balanced, such as they are in a liquid pressure transmitting medium, the parameter n in equation (10) is equal to 3.0. All strains are due to hydrostatic stresses. The assumption of hydrostatic compression led Ahrens and Katz (1962) to use an expression as Cook's method identical to equation (12) with n = 3. If the deformation of the specimen is piston-like, i.e., the side walls are rigid and only the thickness changes, then the value of n is 1.0. If the sidewalls - as in our experiments - are rigid or easier deformable than the buffers in axial direction, $n \le 1.0$.

We found n = 0.622 for run 3.27.

A third possibility to determine density as a function of pressure is an iterative numerical approach. The calculation of the adiabatic bulk modulus at the first pressure step starts with the assumption $\rho = \rho_{\theta}$, i.e., the density do not change within this small pressure-interval. The resulting compressibility is used to calculate the increased density at this pressure, which is used for the next calculation cycle. The result is very close to the data determined by using the EoS published by Ahrens and Katz (1962) and Birch, (1986).

The in situ density evaluation was performed while using unit cell parameters of NaCl derived by XRD.

 K_0 - and K'_T - values published by Birch (1986) were used to calculate the isothermal bulk modulus K_T and the corresponding compressibility κ_T using equation (2) and (4). The V/V_0 - values published by Bridgman (1940) were also used to calculate the isothermal compressibility. Both values agree very well (Figure 14). The difference between the adiabatic (K_s) and isothermal (K_T) bulk moduli is

$$K_S = K_T (1 + \alpha \gamma T) \tag{18}$$

and $\alpha \gamma T \approx 0.01$ at room temperature (Kung and Rigden, 1999) was taken into account.

The detailed comparison of the data showed minor differences. The ultrasonic curves cross the static compression graphs twice. At ≈ 1.2 GPa the compressibility graph derived from the EoS-fitted ultrasonic data intersects the static compression graph first. At ambient conditions the static compressibility is 7 % higher than the dynamical compressibility derived from ultrasonic measurements. This seems to be the result of nonintrinsic compression, e.g., due to a closure of microcracks at the early compression stage in static compression experiments.

Between 2 GPa and 4 GPa the graphs are nearly parallel with up to 3 % higher compressibility derived from ultrasonic measurements. The high pressure intersection is located at 5.3 GPa. At higher pressures the difference seems to increase. At our maximum pressure of 7.71 GPa the static compressibility is again 6.6 % higher than the presented value. This may lead to significant errors for the pressure standard at higher pressure.

In terms of pressure measurement the compressibility calculated from ultrasonic data indicate at 3 GPa about 0.25 GPa higher pressures than derived from static compression data by Bridgman (1940). The ultrasonic data are related to Decker (1971) pressure scale.

The graphs for all calculated NaCl compressibilities, i.e., derived from static compression and from ultrasonic measurements using the equation of state, the empirical deformation model and the numerical approach, were polynomial fitted up to the power of 5, which is required for the range between 1 GPa and 5 GPa. Table 1 presents the coefficients of this fits according to equation (19).

$$K_{r} = A + B_{1}P + B_{2}P^{2} + B_{3}P^{3} + B_{4}P^{4} + B_{5}P^{5}$$
(19)

Figure 15 shows the relation between the Decker (1971) pressure scale and the pressure derived from the ultrasonic measurements of this study using the Eos by Birch (1986). The data were also polynomial fitted up to the power of 5:

$$p_{us} = 0.34611 + 0.6807 p_{De} + 0.01921 p_{De}^{2} + 0.00246 p_{De}^{3} + 8,4777*10^{-4} p_{De}^{4}$$
(20)
+ 5.75971*10⁻⁵ p_{De}^{5}

with p_{us} - pressure derived from ultrasonic measurements of this study

p_{De} - pressure according to Decker (1971)

4. CONCLUSIONS

The results demonstrate the ability to measure the pressure inside of multi-anvil pressure cells standard-free by ultrasonic interferometry. The synchrotron radiation is used to measure the pressure by XRD-techniques using EoS after Decker (1971). The synchrotron radiation is also used for precise in situ sample length and density determination required for the ultrasonic method. Different ways of density determinations were used (using the EoS for NaCl, published by Birch (1986), analyzing the deformation (Ahrens and Katz, 1962), and using an iterative numerical approach) and agreed within <0.1%. Ultrasonic pressure measurement will probably not substitute the XRD-determination completely, because of its higher technical expense, but might be important for a calibrant-free pressure scale determined at very high pressures. However, it seems to become a standard high-pressure method to determine elastic properties of polycrystalline samples parallel to the growing amount and quality of ultrasonic measurements on single crystals under experimental simulated Earth's mantle conditions.

Acknowledgements. We would like to express our special thanks to the editors J. Chen, Y. Wang, T. Duffy, G. Shen, and L. Dobrzhinetskaya for their initiative and patience. We are especially grateful to S. Ganschow for the Ce:YAG crystal, manufactured and put at our disposal by IKZ and J. Kulesza (ADC) for his untiring support. The authors thank M. Kreplin and G. Berger for fabricating tools and cell parts, C. Karger for her software contribution, and H. Witzki and W. Steiner for many initiatives and important technical contributions as well as all colleagues of the high pressure mechanical workshop for their dedicated support.

REFERENCES

Ahrens, T.J., Katz, S., 1962. An ultrasonic interferometer for high-pressure research. J. *Geophys. Res.* 67, 2935-2944.

Birch, F., 1978. Finite strain isotherm and velocities for single-crystal and polycrystalline NaCl at high pressures and 300°K. J. *Geophys. Res.* 83, 1257-1268.

Birch, F., 1986. Equation of state and thermodynamic parameters of NaCl to 300 kbar in the high-temperature domain. *J. Geophys. Res.* 91, 4949-4954.

Boehler, R., Kennedy, G.C., 1980. Equation of state of sodium chloride up to 32 kbar and 500°C. *J. Phys. Chem. Solids* 41, 517-523.

Bridgman, P.W., 1940. Compressions to 50,000 kg/cm². *Physical Review* 57, 237-239.

Brown, J.M. (1999): NaCl pressure standard. J. Appl. Phys. 86, 5801-5808.

Chen, J., Parise, J.B., Li, R., Weidner, D.J., Vaughan, M., 2000. The imaging plate system interfaced to the large-volume press at beamline X17B1 of the national

synchrotron Light Source. In: Manghnani, M.H., Yagi, T. (Eds.), Properties of Earth and planetary materials at high pressure and temperature. *Geophysical Monograph 101*, AGU, Washington, DC, pp. 139-144.

Cook, R.K., 1957. Variation of elastic constants and static strains with hydrostatic pressure: a method for calculation from ultrasonic measurements. *J. Acoust. Soc. Am. 29*, 445-449.

Decker, D.L., 1971. High-pressure equation of state for NaCl, KCl, and CsCl. J. Appl. Phys. 42, 3239-3244.

Frankel, J., Rich, F.J., Homan, C.G., 1976. Acoustic velocities in polycrystalline NaCl at 300°K measured at static pressures from 25 to 270 kbar. *J. Geophys. Res.* 81, 6357-6363.

Fritz, J.N., Marsh, S.P., Carter, W.J., Mcqueen, R.G., 1971. The Hugonoit equation of state of sodium chloride in the sodium chloride structure. In: Lloyd, E.C. (Ed.), Accurate characterization of the high pressure environment. *NBS Spec. Publ. 326*, pp. 201-208.
Holland, K.G., Ahrens, T.J., 1998. Properties of LiF and Al 2 O 3 to 240 GPa for metal shock temperature measurements. In: Manghnani, M.H., Yagi, T. (Eds.), Properties of Earth and planetary materials at high pressure and temperature. *Geophysical Monograph 101*, AGU, Washington, DC, pp. 335-343.

Getting, I.C., 1998. The practical pressure scale: Fixing fixed points and future prospects. *Eos 79*, F830.

Katz, S., Ahrens, T.J., 1963. Ultrasonic measurements of elastic properties of small specimens at high pressure. *High Pressure Meas. Pap. 1962*, 246-261.

Knoche, R., Webb, S.L., Rubie, D.C., 1997. Experimental determination of acoustic wave velocities at Earth mantle conditions using a multianvil press. *Phys. Chem. Earth,*

22, 125-130.

Knoche, R., Webb, S.L., Rubie, D. C., 1998. Measurements of acoustic wave velocities at P-T conditions of the Earth's mantle. In: Manghnani, M.H., Yagi, T. (Eds.) Properties of Earth and planetary materials at high pressure and temperature. *Geophys. Mon. 101*, AGU, Washington, DC, pp. 119-128.

Kung, J., Rigden, S., 1999. Oxide perovskites: pressure derivatives of the bulk and shear moduli. *Phys. Chem. Minerals 26*, 234-241.

Kung, J., Gwanmesia, G.D., Liu, J., Li, B., Liebermann, R.C., 2000. PV3T Experiments: Simultaneous measurement of sound velocities (Vp and Vs) and sample volume (V) of polycrystalline specimens of mantle minerals at high pressures (P) and temperatures (T). *Eos 81*, No. 48, F1151.

Kung, J., Angel, R.J., Ross, N.L., 2001a. Elasticity of CaSnO₃ perovskite. *Phys. Chem. Minerals* 28, 35-43.

Kung, J., Weidner, D.J., Li, B., Liebermann, R.C., 2001b. Determination of the elastic properties at high pressure without pressure scale. *Eos* 82, F1383.

Li, B., Chen, G., Gwanmesia, G.D., Liebermann, R.C., 1998. Sound velocity measurements at mantle transition zone conditions of pressure and temperature using ultrasonic interferometry in a multianvil apparatus. In: Manghnani, M.H., Yagi, T. (Eds.) Properties of Earth and planetary materials at high pressure and temperature. *Geophys. Mon. 101*, AGU, Washington, DC, pp. 41-61.

Li, B., Vaughan, M.T., Kung, J., Weidner, D.J., 2001. Direct length measurement using X-radiography for the determination of acoustic velocities at high pressure and temperature. *NSLS Activity Report*, 2-103-106.

Luth, W.L., 1993. Measurement and control of intensive parameters in experiments at

high pressure in solid-media apparatus. In: Luth, R.W., (Ed.), Experiments at high pressure and applications to the Earth's mantle, Min. Ass. Canada, Short course handbook 21, Edmonton, Alberta, pp. 15-37.

Mao, H.K., Xu, J., Bell, P.M., 1986. Calibration of the ruby pressure gauge to 800 kbar under quasi-hydrostatic conditions. *J. Geophys. Res. 91*, No.B5, 4673-4676.

McSkimin, H.J., 1950. Ultrasonic measurement techniques applicable to small solid specimens. J. Acoust. Soc. Am. 22, 413-418.

Mueller, H.J., Lauterjung, J., Schilling, F.R., Lathe, C., Nover, G., 2002. Symmetric and asymmetric interferometric method for ultrasonic compressional and shear wave velocity measurements in piston-cylinder and multi-anvil high-pressure apparatus. *Eur. J. Mineral.* 14, 581-589.

Mueller, H.J., Schilling, F.R., Lauterjung, J., Lathe, C., 2003. A standard free pressure calibration using simultaneous XRD and elastic property measurements in a multi-anvil device. *Eur. J. Mineral.* 15, 865-873.

Piermarini, G.J., Block, S., Barnet, J.D., Forman, R.A., 1975. Calibration of the pressure dependence of the R₁ ruby fluorescence line to 195 kbar. *J. Appl. Phys. 46*, 2774-2780. Ruoff, A.L., Lincoln, R.C., Chen, Y.C., 1973. A new method of absolute high pressure determination. *J. Phys. D: Appl. Phys. 6*, 1295-1306.

Shen, A.H., Reichmann, H.-J., Chen, G., Angel, R.J., Bassett, W.A., Spetzler, H., 1998. GHz ultrasonic interferometry in a diamond anvil cell: P-wave velocities in periclase to 4.4 GPa and 207°C. In: Manghnani, M.H., Yagi, T. (Eds.) Properties of Earth and planetary materials at high pressure and temperature. *Geophys. Mon. 101*, AGU, Washington, DC, pp. 71-77.

Shimomura, O., Yamaoka, S., Yagi, T., Wakutsuki, M., Tsuji, K., Kawamura, H.,

Hamaya, N., Fukunaga, O., Aoki, K., Akimoto, S., 1985. Multi-anvil type X-ray system for synchrotron radiation. In: Minomura, S., (Ed.), Solid state physics under pressure. Terra Sci. Pub. Comp., 351-356.

Vaughan, M.T., 1993. In situ X-ray diffraction using synchrotron radiation at high P and T in a multi-anvil device. In: Luth, R.W., (Ed.), Experiments at high pressure and applications to the Earth's mantle, Min. Ass. Canada, Short course handbook 21, Edmonton, Alberta, pp. 95-130.

Yagi, T., 1988. MAX80: Large-volume high-pressure apparatus combined with synchrotron radiation. *Eos 69*, 12, 18-27.

Zha, C., Duffy, T.S., Downs, R.T., Mao, H., Hemley, R.J., 1998. Brillouin scattering and X-ray diffraction of San Carlos olivine: direct pressure determination to 32 GPa. *Earth Plan. Sci. Lett. 159*, 25-33.

Zinn, P, Hinze, E., Lauterjung, J., Wirth, R., 1997. Kinetic and microstructural studies of the quartz-coesite phase transition. *Phys. Chem. Earth* 22, 105-111.

Tables

А

 B_1

 B_2

 B_3

 \mathbf{B}_4

 B_5

ultrasonic interferometry and static compression.						
	static	ultrasonics,	ultrasonics,	ultrasonics,		
	compression	this work,	this work,	this work,		
	(Birch, 1986	density from	density from	density from		
		EoS	deformation	numerical		

(Birch, 1986)

0,03907

-0,00523

6.51178*10-4

-1.09931*10-4

2.19162*10-5

-3.17726*10-6

0,04191

-0,0088

0,0018

 -2.43008_*10^{-4}

 1.93729_*10^{-5}

-8.72196*10-7

(Ahrens and

Katz, 1962)

0,03907

-0,00514

6.50735*10-4

 -1.02209_*10^{-4}

1.86429*10-5

 $-2.56361*10^{-6}$

approach

0,03907

-0,00514

6.00421*10-4

 $-9.2428_{*}10^{-5}$

1.83552*10-5

-2.71682*10-6

TABLE 1. Polynomial fit coefficients for the compressibility of NaCl measured by

Figures

Figure 1. DIA-type multi-anvil apparatus MAX80 with Ge solid state detector. The load frames are assembled at a 250 tons hydraulic ram. The Ge solid state detector is also assembled at the press frame and follows the adjustment of the whole apparatus in relation to the X-ray beam.

Figure 2. Load frame, anvils and sample arrangement of MAX80.

(a) Apparatus opened for sample exchange and

(b) vertical cross section with transducer installation at the top anvil.

The tungsten carbide anvils with a steel made keep ring are assembled to bolsters and reaction bolsters, respectively. Driven by the vertical movement of the hydraulic cylinder the load frame and the reaction bolsters generate the movement of the lateral anvils.

Figure 3. Ultrasonic high-pressure set-ups for anvil truncations of 6 mm and 3.5 mm. The smaller set-up was not equipped with a heater to keep the sample cross-section as big as possible, because the strength of reflected ultrasonic waves is a function of the sample diameter.

Figure 4. Transducer arrangements on the rear side of MAX80 anvils.

- (a) Two transducer couples for asymmetrical set-ups
- (b) Single transducer for symmetrical set-up

Figure 5. Electronic equipment for ultrasonic interferometry at MAX80. Rectangular pulses made by an arbitrary waveform generator gate a signal generator resulting in rf pulses or double-pulses. A directional bridge prevents the power burst from hitting the sensitive pre-amplifier. The oscilloscope displays and digitizes the received ultrasonic signals. Amplifiers and an integrated trigger source (5900PR) were used. Transducers were installed on modified top and bottom anvils of MAX80. A computer controlled switch selects the active transducer or transducer pair.

Figure 6. Typical resultant travel time determination using interference pattern. Travel time curves are plotted as a function of frequency at 7.71 GPa. Each point represents a frequency for constructive or destructive interference, and hence can be considered as an independent travel-time determination. The symbols fitted by the horizontal line represent the revealed travel-time. The upper and lower curves represent neighboring fringes of interference pattern.

Figure 7. Determination of sample length under in situ conditions by XRD-scanning. The position of the interface is calculated as half the distance between the last and first appearance of pure XRD-spectra.

Figure 8. High-precision 4-blade slits system

Figure 9. Scheme of X-radiography at MAX80

Figure 10. X-radiography system without camera inside PB-shielding below the XRD-detector.

Figure 11. X-ray shadow graph (a) and its evaluation by density analysis in linear (b) and logarithmic scale (c)

Figure 12. Prefabricated gasket insets prior and after the high pressure run.

Figure 13. Elastic wave velocities v_p and v_s of polycrystalline NaCl at high pressure. Runs 2.2 and 3.10 use 8 mm set-ups for 6 mm anvil truncation; run 3.27 uses a 5.5 mm set-up for 3.5 mm anvil truncation.

Figure 14. Compressibility of NaCl measured by ultrasonic interferometry and static compression: The calculation of compressibility from elastic wave velocities require the density as a function of pressure. The in situ density was determined by analysing the sample deformation (deformation fit), using published EoS (EoS fit) and successive approximation. The X-axis is related to the Decker (1971) pressure scale.

Figure 15. Pressure measured by ultrasonic interferometry in this study vs. Decker (1971) pressure scale related to the EoS by Birch (1986).

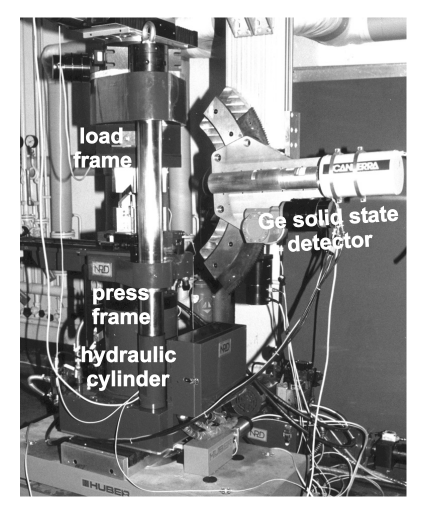
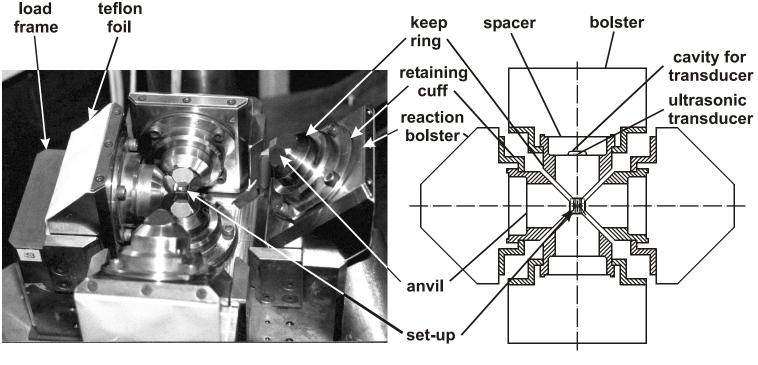


Figure 1. DIA-type multi-anvil apparatus MAX80 with Ge solid state detector. The load frames are assembled at a 250 tons hydraulic ram The Ge solid state detector is also assembled at the press frame and follows the adjustment of the whole apparatus in relation to the X-ray beam



(a)

(b)

Figure 2. Load frame, anvils and sample arrangement of MAX80.

(a) Apparatus opened for sample exchange and

(b) vertical cross section with transducer installation at the top anvil.

The tungsten carbide anvils with a steel made keep ring are assembled to bolsters and reaction bolsters, respectively. Driven by the vertical movement of the hydraulic cylinder the load frame and the reaction bolsters generate the movement of the lateral anvils.

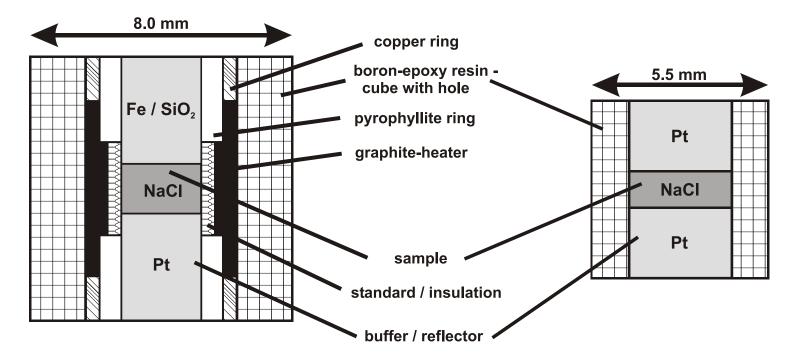
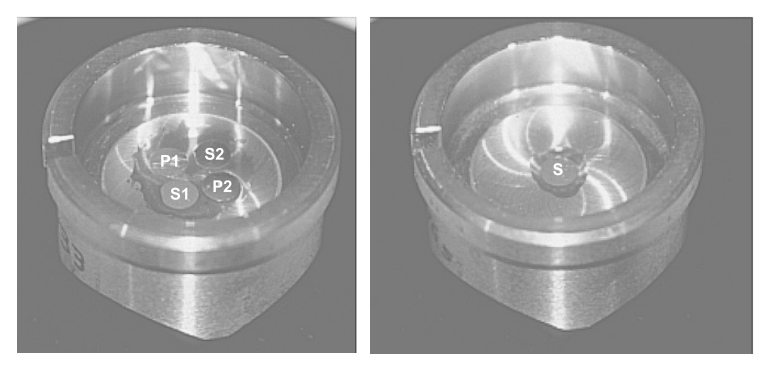


Figure 3. Ultrasonic high-pressure set-ups for anvil truncations of 6 mm and 3.5 mm The smaller set-up was not equipped with a heater to keep the sample cross-section as big as possible, because the strength of reflected ultrasonic waves is a function of the sample diameter.



(a)

(b)

Figure 4. Transducer arrangements on the rear side of MAX80 anvils. (a) Two transducer couples for asymmetrical set-ups (b) Single transducer for symmetrical set-up

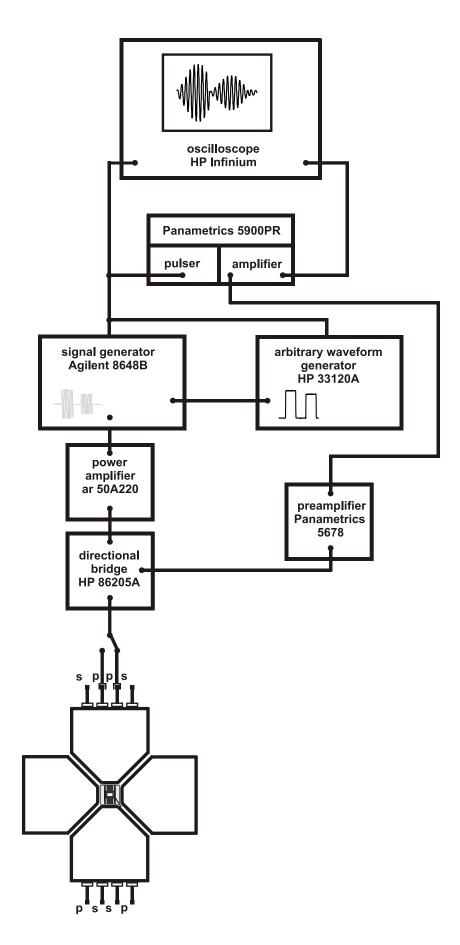


Figure 5. Electronic equipment for ultrasonic interferometry at MAX80. Rectangular pulses made by an arbitrary waveform generator gate a signal generator resulting in rf pulses or double-pulses. A directional bridge prevents the power burst fromhitting the sensitive pre-amplifier. The oscilloscope displays and digitizes the received ultrasonic signals. Amplifiers and an integrated trigger source (5900PR) were used. Transducers were installed on modified top and bottom anvils of MAX80. A computer controlled switch selects the active transducer or transducer pair.

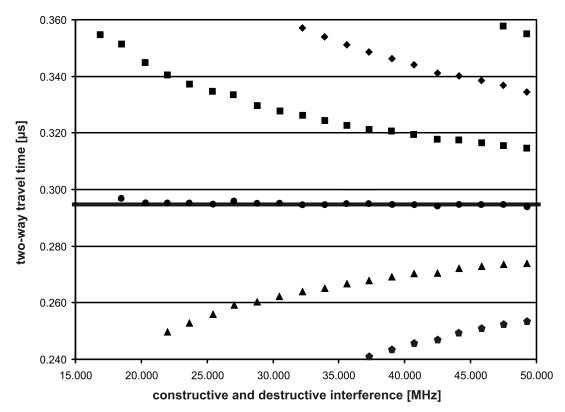


Figure 6. Typical resultant travel time determination using interference pattern. Travel time curves are plotted as a function of frequency at 7.71 GPa. Each point represents a frequency for constructive or destructive interference, and hence can be considered as an independent traveltime determination. The symbols fitted by the horizontal line represent the revealed travel-time. The upper and lower curves represent neighboring fringes of interference pattern.

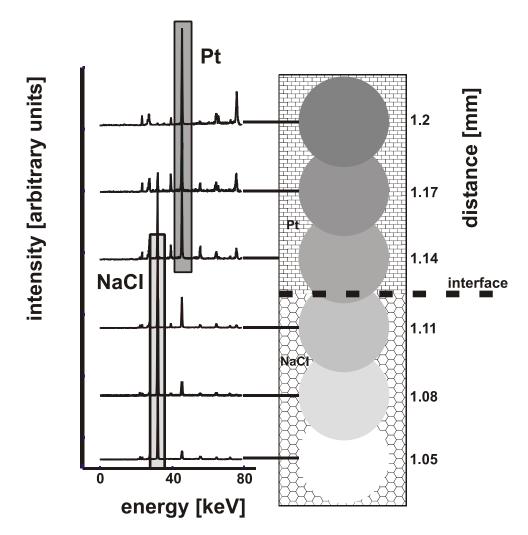


Figure 7. Determination of sample length under in situ conditions by XRD-scanning. The position of the interface is calculated as half the distance between the last and first appearance of pure XRD-spectra.

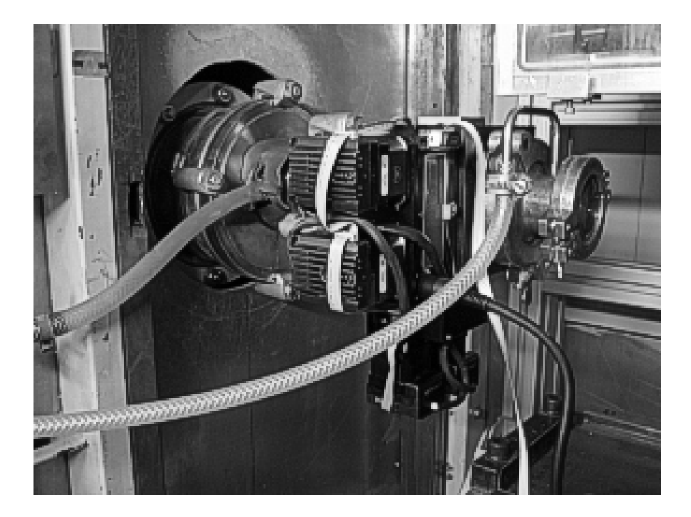


Figure 8. High-precision 4-blade slits system

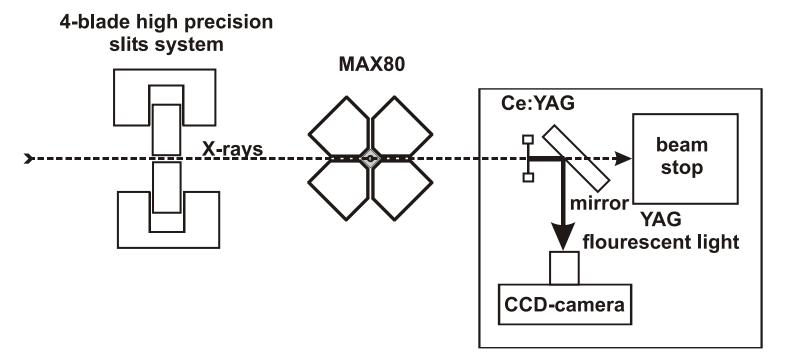


Figure 9. Scheme of X-radiography at MAX80

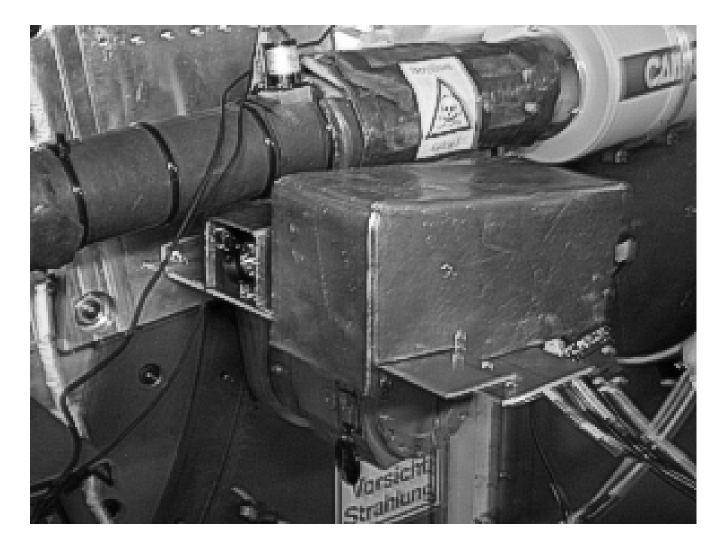


Figure 10. X-radiography system without camera inside PB-shielding below the XRD-detector

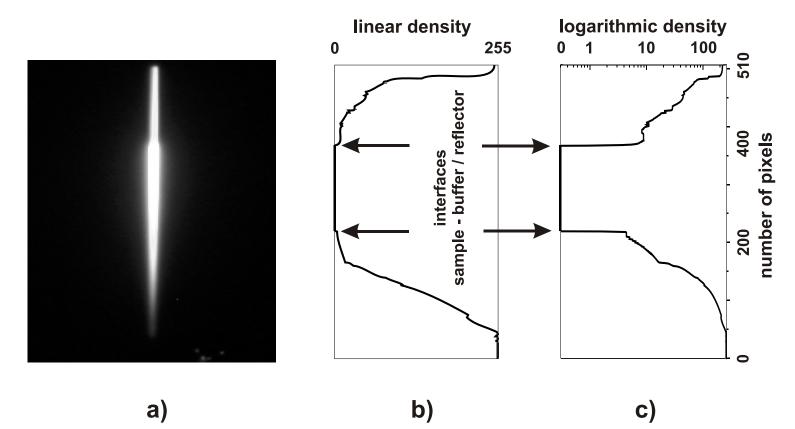


Figure 11. X-ray shadow graph (a) and its evaluation by density analysis in linear (b) and logarithmic scale (c)

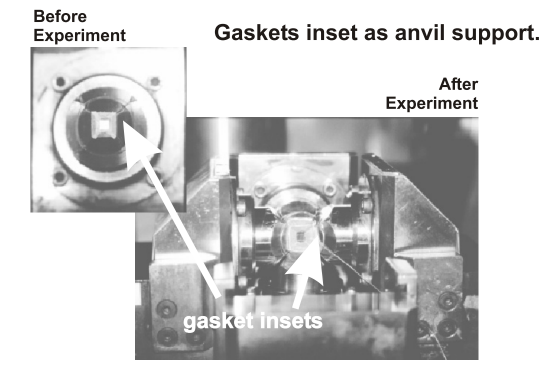


Figure 12. Prefabricated gasket insets prior and after the high pressure run.

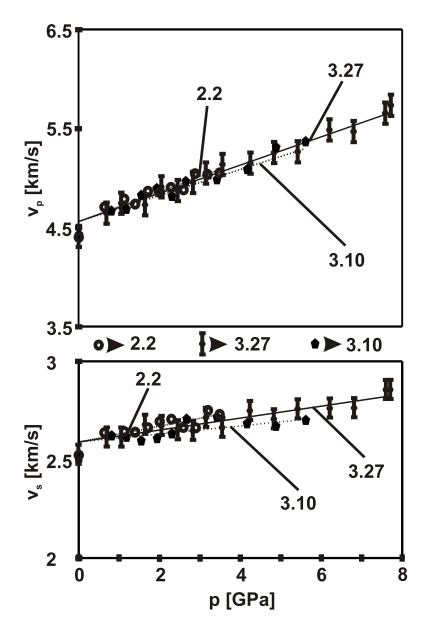


Figure 13. Elastic wave velocities v_p and v_s of polycrystalline NaCl at high pressure. Runs 2.2 and 3.10 use 8 mmset-ups for 6 mmanvil truncation; run 3.27 uses a 5.5 mmset-up for 3.5 mm anvil truncation.

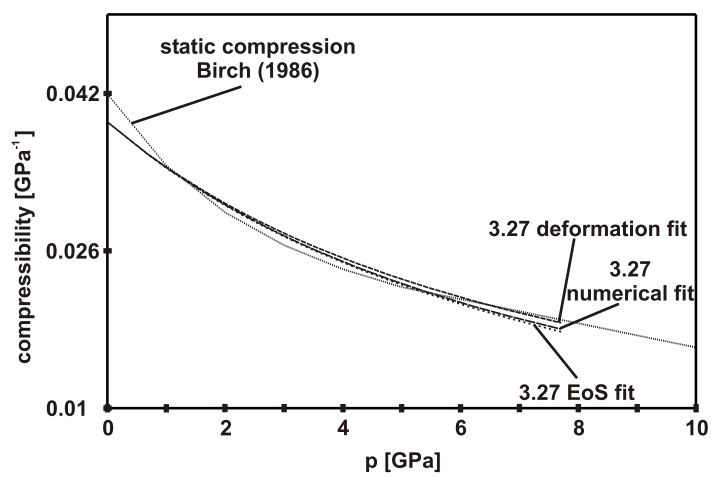


Figure 14. Compressibility of NaCl measured by ultrasonic interferometry and static compression: The calculation of compressibility from elastic wave velocities require the density as a function of pressure. The in situ density was determined by analysing the sample deformation (deformation fit), using published EoS (EoS fit) and successive approximation. The X-axis is related to the Decker (1971) pressure scale.

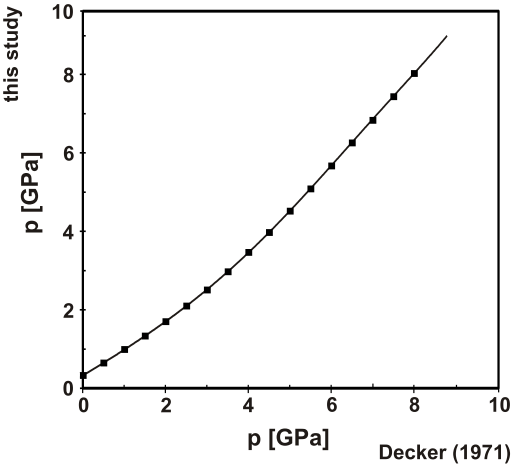


Figure 15. Pressure measured by ultrasonic interferometry in this study vs. Decker (1971) pressure scale related to the EoS by Birch (1986).