High Pressure Techniques

M. Guthrie, Geophysical Laboratory
High pressure: a tool for material science

- Without varying external parameters, chances of understanding materials is rather limited (e.g. SC...).
- Popular variables are T,X (less common are H,E,Φ...)

P is an underused but uniquely powerful variable
High pressure: a tool for material science

• P is essential to explore full versatility of inter-atomic bonds

Vital to synthesise diamond

• Contemplate new spectrum of sp$^3$ C allotropes e.g. mesoporous diamond, amorphous diamond, hydrogenated carbons...
High pressure: a tool for material science

- Can directly modify electronic properties (subtle)

Can tune superconducting transition $T_c$

[X.-J. Chen et al. Nature 466 950 (2010)]
High pressure: a tool for material science

- Can directly modify electronic properties (dramatic)

**Transparent sodium**
(at 199 GPa)

**Metallic oxygen**
(at 133 GPa)
[G. Weck et al PRL 102 255503 (2009)]

Metal ↔ Insulator transitions
High pressure: a tool for material science

• P can provide direct access to metastable states

• Pressure induced amorphisation (ice, SiO$_2$, ZrW$_2$O$_8$...)

• Pressure induced Crystallisation (ice, Si, Ge, BMG...)

• Pressure amorphous-amorphous transitions (ice, SiO$_2$, GeO$_2$, Si...)

![Graphical representation of high pressure effects on materials]
Pressure can also denature proteins...

On the physics of pressure denaturation of proteins

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Abstract

We show that the entropic effect originating from the translational movement of water molecules plays critical roles in the pressure-induced denaturation of proteins. In our statistical-mechanical method, the partial molar volume

Also claims (controversial) that some bacteria can survive extreme pressures (in excess of 1.6 GPa)

Microbial Activity at Gigapascal Pressures

Anurag Sharma,* James H. Scott,* George D. Cody, Marilyn L. Fogel, Robert M. Hazen, Russell J. Hemley, Wesley T. Huntress

We observed physiological and metabolic activity of *Shewanella oneidensis* strain MR1 and *Escherichia coli* strain MG1655 at pressures of 68 to 1680 megapascals (MPa) in diamond anvil cells. We measured biological formate oxidation at high pressures (68 to 1060 MPa). At pressures of 1200 to 1600 MPa, living bacteria resided in fluid inclusions in ice-VI crystals and continued to be viable upon subsequent release to ambient pressures (0.1 MPa). Evidence of microbial viability and activity at these extreme pressures expands by an order of magnitude the range of conditions representing the habitable zone in the solar system.
Pressure scale?

SI unit for pressure: Pascal, Pa (1 Nm\(^{-2}\))
i.e. Force/Area

Research at neutron and x-ray facilities is routinely conducted at pressures measured in GigaPascals, GPa*.

Reference
Atmospheric pressure ~ 0.0001 GPa
Deepest point of the ocean ~ 0.1 GPa
Stability field of diamond > 5 GPa
Center of the Earth: ~350 GPa

(*I may slip into kbar = 1000 bar during talk…conversion is easy 1 GPa = 10 kbar)
How do you generate high pressures in the lab?

Mechanical compression of gases possible since early in the industrial revolution. Gas pressures up to ~200 bar (0.02 GPa) are common.

200-300 kPa (2-3 bar) 1.5 MPa (15 bar) 20 MPa (200 bar)

Higher gas pressures of up to ~0.5 GPa in oil & gas industry

Compressing solids and liquids is much harder, and was considered impossible until early 20th century.

What’s the difference between compressing a gas and compressing a solid?
How do you generate high pressures in the lab?

Wide range of gas compressors (see e.g. http://en.wikipedia.org/wiki/Gas_compressor)

For highest gas pressures - one dominant technique: the piston cylinder.

\[ P = \frac{F}{A} \]

- Pressure, \( P_1 \) applied to Area, \( A_1 \)
- This generates force, \( F = P_1 \times A_1 \)
- This force is applied to smaller area, \( A_2 \)
- Generating pressure \( P_2 = \frac{F}{A_2} = P_1 \times \frac{A_1}{A_2} \)

The greater the pressure, the simpler the device
Going beyond the piston-cylinder

How about solids? Can they be compressed using a piston-cylinder?

Yes…Maximum pressures of ~ 2 GPa are relatively routine (max ~ 5 GPa)…this is already enough to compress some solids (e.g. ~15 ice phases below 3 GPa – by rearranging molecules)

But a radically different design was required to go to higher pressure.

This came courtesy of Percy Bridgman in the early 1900’s (and subsequently earned him a Nobel Prize)

“You, Mr. Bridgman, have succeeded in doing what was once considered impossible. By the use of new alloys and by other ingenious devices you have been able figuratively speaking, to bring into your laboratory parts of the interior of the earth or of other places where no human being is able to exist, and you have been able there to examine the physical and chemical properties of a quantity of different substances under the enormous pressures you have created. You have thus been able to reveal a number of strange phenomena in the behaviour of matter under other circumstances than those which we consider to be normal.”

P. W. Bridgman
1882-1961

— Sigurd Curman, President Royal Academy of Sciences, Prior to presenting Bridgman’s Nobel Prize in physics 1946
“Stuck between a hard place and a hard place”

Bridgman’s insight was a technique based around an opposed anvil design – with it he eventually reached ~40 GPa

Three elements of the opposed anvil technique:
1) Two anvils made of a hard material
2) A force to push these anvils together
3) A gasket made of a material that is strong, but able to flow

These same principles apply to the majority of high-pressure cells operating today above ~2 GPa at synchrotron and neutron sources.
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What is a hard material? Bridgman used a composite of WC and cobalt. Other materials used are pure WC, sapphire ($\text{Al}_2\text{O}_3$), moissanite (SiC), c-BN…

But in almost all cases, the best material is diamond.

Diamond anvils are either single-crystal or poly-crystalline. PCD available (sintered, typically with Co binder). Also in last 10 years ultra-hard nano-PCD (HIME-DIA)

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The amount of force (and how it’s applied) depends on the area of the sample and the required pressure.

X-ray cells (<1 tonne) (screw, membrane, piezo actuator)

“conventional” neutron cells 150-500 tonnes (hydraulic presses)

Multi-anvil 1000-6000 tonnes

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1) Gasket (typically metal, but can also be composite material)
2) Apply force to ‘indent’ gasket:
   • Work hardens gasket
   • Forms support for diamond tips
   • Stable geometry (thin)
3) Drill hole for sample
   (for DAC’s need EDM or laser as hole is very small)
4) Load sample, pressure calibrant* and pressure medium*

(* discussed soon)
“Stuck between a hard place and a hard place”

Seal cell by applying further force. As gasket can flow, it follows pressure gradient, moving away from sample.

In process, thinning and reducing volume available to sample – increasing pressure.

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Pressure measurement

As with any experiment, accurate knowledge of the variable you control is very important.

Pressure is measured the same way any other variable is:
**Calibrate something with a physical response to variable of interest**

Example 1) **Ruby fluorescence.**
- Probably the most ubiquitous pressure sensor above 2 GPa
- Under laser light, ruby fluoresces with particular spectrum
- The wavelength (colour) shifts in a known way with pressure

Example 2) Known equation-of-state of calibrant
- If ruby isn’t an option (opaque anvils, high temperature, reactivity)
- Can load a secondary sample with a known pressure-volume relation. Use diffraction to determine volume – and, therefore, pressure.

Others… raman shift of C$^{13}$, pressure-load curves, ...

**How are the calibrants calibrated?**

Typically shock wave data (discussed later) can give a direct equation of state.
Pressure media

Imagine hard sample directly squeezed between two diamonds...

Results in enormous strain (often many GPa)

Solution is to surround the sample with a medium that is very soft...

• Because medium is soft, it can’t sustain a P gradient
• Sample feels equal pressure on all sides
• Fragile single-crystals, bio-samples can be compressed
• Best media are the inert gases: He, Ne, Ar
• Methanol:ethanol, silicon oil, fluorinert also used
(Need medium that doesn’t react with sample)
Beyond two opposed anvils

For large volumes, an alternative technique uses multiple (typically 6-8) anvils. Well suited for liquid/glass diffraction studies, tomography, element partitioning studies...

Elements:
• Usually uniaxial force (from very large capacity press (+1000 tonnes) but 6 axis presses exist
• 6 anvils with square faces come together to form a cubic sample volume
• 8 anvils – cubes with corner cut off - form octahedral sample space. This assembly can be pressed inside 6 regular anvils (double-stage design)

Sample space is typically filled with:
• Gasket
• Thermal insulation
• Graphite Heater
• Contacts for thermocouples/heater
• Pressure medium/sample encapsulation

2-stage design with PCD anvils, can reach ~80 GPa

Reference: http://www.gps.caltech.edu/~jed/Multianvilpage.html; http://www.misasa.okayama-u.ac.jp/~hacto/facility_e.html
Dynamic compression

Completely different route to achieve highest pressures is via dynamic techniques:

Shockwaves can generate exceptionally high P & T over short time period:
• Nuclear
• Gas gun
• Lasers (NIF)

Under shock, samples experience conditions that lie on a locus in PT space called “Hugoniot”

At NIF expect to reach TPa and $10^4 \text{ K}$ regime (Centre of Jupiter)

Alternative techniques using Piezo actuators can look at dynamic phenomena.
Science at high pressure

Have looked at ways to generate, control and measure extremely high pressures. In order to conduct science, need way to probe effect of pressure on sample material.

Great variety of probes:

**Visual observations**
Phase transitions (solid-solid, melting, conductivity), single-crystal growth

**Laser-based**
Raman, UV & IR spectroscopy
Brillouin

**Transport measurements**
Electrical conductivity, magnetic susceptibility

**Others…**
sound velocity, DTA...

**Focus here on synchrotron x-ray and neutron based probes**

**Variety of techniques**
- Above 0.6 GPa, neutrons limited to diffraction, phonon measurements, tomography.
- In contrast, huge (and rapidly expanding) range of synchrotron x-ray techniques: (XRD, XAS, XMCD, XRS, XES, IXS, NRIXS, transmission density, tomography...)
Neutrons have many unique capabilities

1) Scattering length is not linearly dependent on atomic number
- neutron diffraction is a great tool for studying light atoms. It’s the only technique that can precisely locate protons (deuterons), Be, B$^{11}$, C, N, and O are strong scatterers
- possibility of negative scattering lengths (e.g. H) means specific pair correlations can be removed
- isotopic substitution can greatly enhance contrast and can also simplify analysis of non-crystalline matter.

2) Absorption cross sections also not linearly dependent on atomic number:
Li$^6$, H, B$^{10}$ are strong absorbers. Pb is transparent.

3) Neutrons have an intrinsic magnetic moment
- They are scattered by nuclear spins and sensitive to magnetic order.

4) Scattering is via inter-nuclear interactions. Pointlike.
- No atomic form factor, so high Q-vectors are accessible.
Leading to exceptional real-space resolution.

[Neutron imaging and applications, Bilheux et al]
Neutrons have one major disadvantage

Photons from 60W bulb!

This is a challenge for high pressure because samples are small…
Examples of High-Pressure Neutron science

**Broken symmetry in hydrogen**

**Salty Ice VII**

**Magnetic ordering in solid O$_2$**

**Stability of methane hydrate**

**Phonon dispersion in ice I$_h$**

**Structure of liquid ammonia**
Guthrie et al PRB(2012).
Neutrons science at high pressure

Mature neutron facilities with HP programmes:

Europe

- ISIS, UK
- ILL, France
- Saclay, France
- Dubna, Russia
- SINQ, Switzerland

US

- LUJAN CENTER, LANL
- HFIR, ORNL
- McClellan, UC Davis
- IPNS, ANL (closed)

• Typical max P ~ 25 GPa
• Max T ~ 1500 K
• Min T ~ 4 K
Neutrons science at high pressure

New neutron facilities with HP programmes:

- JPARC, Japan
- IPNS, ANL (closed)
- SNS, Oak Ridge

- Typical max pressure ~ ? GPa
- Max temperature ~ ? K
- Min temperature ~? K
SNAP – high pressure at the SNS

• Highly pixelated area detectors (Anger cameras) give simultaneous access to large volumes of reciprocal space.
• Movable detectors mean wavelength coverage can be swept from low to high Q-vectors (or high to low d-spacing)
• Moveable flight tube can be replaced with different focusing optics (Elliptical or KB).
• Precise (1um) stage permits alignment of very small samples.
• Highly versatile diffractometer: can study single-crystals, powders or even liquid structure
SNAP – high pressure at the SNS

Conventional HP neutron sample volumes are $\sim 100\text{mm}^3$ and require 200-500 tonne presses.

But the intense flux at SNS means samples can be small.

Is it possible to do neutron work with a DAC?

At NXS 2011, showed this dataset from SNAP.

And claimed...

“high-quality diffraction data up to 60 GPa in 1-2 years”
SNAP – high pressure at the SNS

After 2 years of development, have huge improvement in data quality...

First real structural measurements with neutrons at pressures up to 50 GPa...

\[ \chi^2 = 1.49 \]

Neutron diffraction observations of interstitial protons in dense ice

Malcolm Guthrie, Reinhard Boehler, Christopher A. Tulk, Jamie J. Molaison, António M. dos Santos, Kuo Li, and Russell J. Hemley

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Contributed by Russell J. Hemley, May 16, 2013 (sent for review April 22, 2013)

The motif of distinct H2O molecules in H-bonded networks is believed to persist up to the densest molecular phase of ice. At even higher pressures, where the molecule dissociates, it is generally assumed that the proton remains localized within these same networks. We report neutron-diffraction measurements on D2O that reveal the location of the D atoms directly up to 52 GPa, a pressure in the molecular geometry up to a reported 20 GPa (19) (see comment in Materials and Methods). A lack of in situ neutron-diffraction data above this pressure has precluded direct measurement of the D locations on the approach to symmetrization. By developing a supported diamond-anvil cell (S-DAC) (12) and coupling this with the intense radiation flux at the dedicated high-
SNAP – high pressure at the SNS

As of Aug 2013 got to ~94 GPa!

Aiming for >100 GPa by the end of 2013!

This breakthrough is not only about high pressure. Low T, High T, gas loading, in situ spectroscopy,…a more synchrotron-like neutron experience.
PLANET and the “Pressure Princess”

J-PARC Japanese spallation source (design spec of 1MW) is operational
Multi-anvil based HP neutron beamline: PLANET

Bad luck…
Mar 2011 – Fukushima earthquake set back construction
May 2013 – Accident at Hadron Facility resulted in shut-down

Once fully operational will be competitive with SNS
Future Neutron Sources

Construction start 2014
First neutrons <6 years

Begin construction 2020?
“Cold” neutron source

Bright future for high pressure neutron science!
X-ray science at high pressure

Access to high pressure at synchrotron sources has exploded in last 10 years.

All major synchrotrons have dedicated high pressure beamlines (e.g. ESRF, APS, SPring-8, Petra-III, NSLS, NSLS-II (proposed)). Extreme conditions are an integral part of the (ongoing) APS upgrade.

Beyond dedicated beamlines...portable high pressure apparatus are extremely wide-spread.

With few exceptions almost all synchrotron techniques you’ll hear about in NXS2013 can be combined with high pressure.
High-pressure diffraction with x-rays

Modern HP beamlines deliver extremely intense, low divergence beams 2-5μm
Coupled with laser heating – can reach >300 GPa and >3000 K

- Gasket hole ~ 60 μm (culet =150 μm, bevel diameter = 300 μm)
- X-ray beam ~ 6 x 7 μm
- Double-sided Nd:YLF laser heating

Schematics of diamond anvil cell setup at 13ID-D at the Advanced Photon Source
Nano-probe diffraction at high P

Beams orders of magnitude smaller than neutrons permit sub-micron studies (could be route to TPa pressures?)

Using 200 nm focused x-ray beam we can...

L Wang, PNAS (2010)

Observe 20 GPa/µm gradient & peak-pressure in 1-µm area

Separate submicron Pt, Re, Fe samples

Conduct single-crystal XRD on submicron powder
Time-resolved diffraction at high P

Add small “t” to variables

• Continuous exposures
• Rietveld refinement every 7 ms
• Structural information throughout transitions
Shocking Diffraction

Very rapid acquisition (ns) facilitates measurements *during shock wave*.

- Phase transitions
- Deformation processes and fracture dynamics
- Dynamics of chemical reactions

DCS is a facility at the APS to study structure and dynamics in these thermo-mechanical extremes.

MOU signed late 2012 for construction at Sector 35

http://www.dcs-aps.wsu.edu/
X-ray absorption spectroscopy (XAS)

Direct insight into local structure and bonding environment

- Pre-edge position and intensity: oxidation state
- Edge height: concentration
- XAFS: coordination & structure

Coordination change in GeO$_2$ glass measured with XANES

Baldini et al PRB (2010).

Wilke et al, Amer. Min. 2001
High-pressure x-ray emission spectroscopy (XES)

Observations of high spin-low spin transitions in 3d elements

FeO & Fe$_2$O$_3$

(Fe,Mg)O & (Fe,Mg)SiO$_3$

Badro et al, PRL 1999
Badro et al, PRL 2002

Badro et al, Science 2003
Badro et al, Science 2004
Li et al, PNAS 2004
Lin et al, Science 2007
Lin et al, Nature Geo. 2008
Resolution above is \( \sim 1\, \text{um} \). Using TXM techniques 20nm is possible. Also, coherent diffraction imaging has been used to image strain dist. In gold nano-particles.
Combining X-rays and Neutrons

As neutron capabilities increase, there’s great scope for complementary experiments.
Summary

• Pressure is a powerful modifier of the physical properties of matter
• In the lab, we are able to achieve static pressures and temperatures approaching the centre-of-the-Earth (and dynamic pressures approaching centre of Jupiter – DCS)
• Scientific capabilities are ‘technique-driven’, demanding materials with most extreme properties of strength and hardness.
• Synchrotron HP diffraction and XAS techniques are mature, with a huge diversity of x-ray techniques continually being developed.
• Neutrons can make a powerful contribution to HP science, especially in diffraction.
• Now is beginning of new period of growth in neutron capabilities based at new generation of intense sources, such as SNS.
• Combination of x-ray and neutron science will become increasingly important as scope of neutron capabilities improves in next several years
“Although mathematics cannot be avoided...the aim has been to write a book that most scientists will still find approachable. To this end, the first two chapters [provide] a background in the mathematics and physics that are implicitly assumed in other texts. Thereafter, the philosophy has been one of keeping things as simple as possible.”