

LHC Project Note 7

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Synchrotron radiation induced gas desorption from a prototype LHC beam screen at cryogenic temperature

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Summary

The performance of the vacuum system of the super conducting proton storage rings LHC, will depend critically on the synchrotron radiation induced gas desorption and on the re adsorption of molecules on the cold surfaces. The present design of the system is based on a so-called 'beam screen' inserted in the 1.9 K cold bore of the magnets. Gas molecules desorbed will therefore readsorb on the beam screen which is held at a temperature between 5 and 20 K. Pumping slots in the beam screen enable some of the desorbed gas to be pumped onto the 1.9 K surface of the cold bore.

1. Introduction

In the present design of the Large Hadron Collider (LHC)[¹] the beam screen consists of a 1 mm thick, square section (outer dimensions 38 mm x 38 mm) stainless steel tube, with a 50 μ m layer of Cu covering the whole of the inside surface. The beam screen will operate at a temperature of between 5 K and 20 K.

The Cu coated beam screen will be inserted in the 1.9 K cold bore tube of the superconducting magnet and it is this Cu surface which will be subjected to bombardment by the 43.8 eV critical energy synchrotron radiation from the 7.0 TeV protons with the subsequent photon induced gas desorption. Pumping of the system will rely entirely on cryosorption - partly on the cold surfaces of the beam screen and partly through pumping slots in the screen on the surface of the magnet cold bore.

In this note we report on the measurements of the synchrotron radiation induced gas desorption from two prototype beam screens at 77 K and at a temperature between 4.2 and 10 K using synchrotron radiation with a critical energy of 50 eV from the VEPP-2M storage ring at the Budker Institute of Nuclear Physics in Novosibirsk, Russia.

2. Beam Screen Preparation

A schematic cross-section of the proposed beam screen for LHC is shown in Figure 1. The prototype beam screens used in this experiment each consisted of a 1 m long, square section 304L stainless steel tube (38 mm x 38 mm) with a strip of Cu 0.2 mm thick and 18 mm wide bonded to the inside of each flat side. The bonding was carried out by a skive inlay process[2] which essentially entails machining a fresh stainless surface before roll bonding the Cu onto it. In the process the resulting Cu/stainless steel sandwich was reduced in thickness to 1 mm. Two cooling pipes were soft soldered on to the outer surface.

Along the length of the beam screen were four rows of pumping slots 10 mm x 1 mm, separated longitudinally by 5 mm and covering \sim 2% of the internal surface.

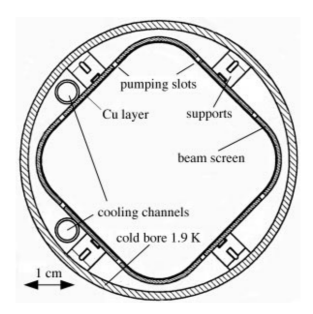


Figure 1: A schematic diagram of the dipole magnet cold bore with the square beam screen containing pumping slots and two cooling channels to maintain the screen between 5 K and 20 K.

The complete beam screen was cleaned by degreasing in an alkaline detergent and then acid etched in a mixture of H_2SO_4 , HNO_3 and HCl (only the Cu was attacked by the acid solution) followed by passivation of the Cu in CrO_3 (chromic acid). After rinsing in demineralized water, the beam screens were dried in a jet of N_2 . The beam screens were transported to Novosibirsk in a sealed plastic bag filled with dry N_2 .

3. Experimental

3.1 Beam line and cryogenic system

Desorption yields were measured on a dedicated synchrotron radiation beam line on the VEPP-2M electron-positron storage ring. This facility has been described fully in previous publications[3,4] but for completeness, and since some modifications had to be made, a brief description of the apparatus will be given here.

Two beam lines, one for high intensity photon irradiation (SSC II) and one for low intensity (SSC I), were originally constructed to measure photon induced gas desorption yields at room temperature and 4.2 K. These operated at a critical energy of 284 eV to provide input data for the design of the cold vacuum system of the 20 TeV Superconducting Super Collider (SSC). On the high intensity beam line, that closer to the storage ring, a cryostat had been installed which enabled measurements to be made at 4.2 K.

The nominal energy of the VEPP 2M storage ring is around 500 MeV but to reproduce the low critical energy of the LHC it was run at an energy of 300 MeV which corresponds to a critical energy of 50 eV, close to the 43.8 eV of the LHC. At this energy the maximum beam current is about 300 mA. The linear photon flux incident on the test chamber is 1.57 x 10¹⁴ photons s⁻¹ m¹ mA⁻¹, which is approximately 12% lower than for LHC with 1.78x10¹⁴ photons s⁻¹ m⁻¹ mA⁻¹. In practice, at 300 MeV the beam lifetime was only some minutes and by reinjecting every few minutes the average beam current was kept above 200 mA. The synchrotron radiation was incident on the corner of the beam screen at a glancing angle of 10 mrad and illuminated almost the complete 1 m length. Total doses of up to 1x10²² photons m⁻¹ could be accumulated over a period of about 7 days. During the last part of this experiment (run #4), the critical energy was increased to 284 eV, the design value of the SSC.

A schematic view of the experimental set-up is shown in Figure 2. In the LHC the beam screen will run at a temperature between 5 K and 20 K and will be inserted in the magnet cold bore which will be at a temperature of 1.9 K. Thus the cryostat shown in Figure 2 was modified to be able to cool the beam screen independently, using either liquid N_2 at 77 K or gaseous He evaporated from a liquid He bath from which the flow was regulated to maintain a constant temperature of 10 K on the outlet of the beam screen cooling circuit. In turn, the beam screen was surrounded by the cryostat which was maintained throughout the experiment at a temperature below 3 K.

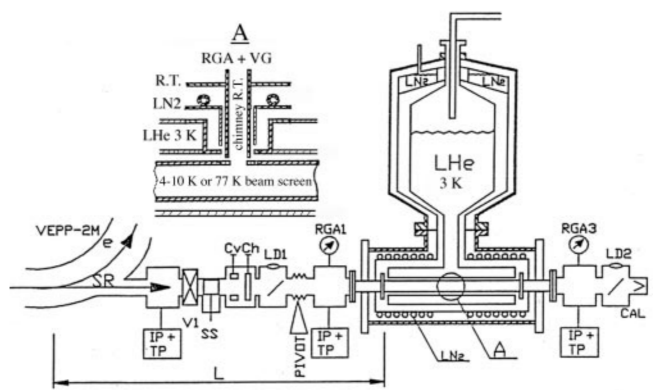


Figure 2: The experimental set-up for the cold beam screen experiment. V1-vacuum valve; Cv, Ch-vertical and horizontal collimators; SS-safety shutter; LD1, LD2;-luminescent screens; VG-ionisation gauge; RGA-residual gas analyser; IP+TP-combination ion and Ti-sublimation pump; L=1.75 m. A calorimeter is mounted at the end of the line. The insert \underline{A} shows a schematic view of the pressure measuring arrangement between the test chamber with the 3 K cold bore and the beam screen which could be adjusted to temperatures between 293 K and 10K. Also shown is the arrangement of the room temperature pressure gauges and the 'chimney' at approximately room temperature leading to the cold test volume.

Of the gases desorbed by the synchrotron radiation, H_2 , CH_4 , CO and CO_2 , only H_2 has a significant saturated vapour pressure in the range 4.2 to 10K (>10⁻⁶ Torr). This H_2 has to be pumped by the cold bore surrounding the beam screen. Thus the cold bore temperature in the experiment should be such that the saturated vapour pressure of the condensed H_2 is less than about 10^{-10} Torr. This implies that its temperature must be less than 3 K. Due to this fact it was not necessary to cool the cold bore to the LHC value of 1.9 K and longer experimental runs could be carried out for a given liquid He capacity.

3.2 Gas density and molecular desorption yield

In the experiment, by necessity, all vacuum gauges and the gas analyser were at room temperature. In an earlier experiment $[^5]$ the chimney connecting these instruments to the cold desorbing zone was at 77 K. It was reasoned that, with this chimney at a higher temperature, but still compatible with heat input to the cold zone, there would be an improved chance of being able to detect all of the desorbed gases, including CO_2 . Thus the cryostat was modified to have the temperature of the chimney close to 293 K and thus avoid any sticking of the molecules once they leave the cold beam screen. The insert in Figure 2 shows schematically the details of the

position of this room temperature chimney with respect to the measuring instruments, the thermal screen at 77 K, the cold bore at <3 K and the beam screen.

The gas density, n, and the pressure, P, inside the beam screen at the temperature T was calculated from the room temperature measurements (subscript RT) using the expressions

$$n = n_{RT} \sqrt{\frac{T_{RT}}{T}}$$
 and $P = P_{RT} \sqrt{\frac{T}{T_{RT}}}$

respectively.

The contribution of the background pressure, Po was measured at many instances throughout the series of experiments by stopping temporarily the photon beam to the test system (as shown by the lower set of measured points in Figure 4). For the evaluation of the molecular desorption yield, h, the background values were subtracted to arrive at the net increase in gas density inside the beam screen

$$\Delta n = \frac{(P - P_o)_{RT}}{k \sqrt{T T_{RT}}}$$

and finally

$$\eta = \frac{\Delta n \ S}{\dot{\Gamma}}.$$

Here k is the Boltzmann constant, S represents the linear pumping speed and ris the linear photon flux in the test chamber as provided by the VEPP-2M storage ring. The pumping by the surface of the beam screen and by the pumping slots in the beam screen is proportional to the mean molecular velocity \bar{v}

$$\overline{v} = \sqrt{\frac{8 k T}{\pi m}} \approx 145 \sqrt{\frac{T}{M}}$$
 (m s⁻¹)

where m is the mass of the molecules and M their molecular weight.

For a sticking probability of molecules, s, the resulting pumping speed per unit length of the beam screen with surface area F is

$$S = \frac{1}{4}\overline{v} \ s \ F.$$

Combining the above expressions gives for the molecular desorption yield

$$\eta = \sqrt{\frac{1}{2\pi}} \frac{s F}{\dot{\Gamma}} \frac{(P - P_o)_{RT}}{\sqrt{m k T_{RT}}}.$$

It is interesting to note, that this result for h becomes independent of the temperature T of the cold system.

Entering the numerical constants, and using units of Pa for the pressure gives
$$\eta \approx 2.64 \cdot 10^{24} \, \frac{s \, F}{\dot{\Gamma}} \, \frac{(P - P_o)_{RT}}{\sqrt{M \, T_{RT}}} \quad \text{(mol/phot)}$$

For the evaluation of the data from the different experimental runs two extreme situations occur: when the screen is at sufficiently high temperature or saturated with gas such that is does not adsorb, s=0; when the screen is at low temperature and not saturated, 0 < s < 1. In all situations a value of s=1 has been assumed for the area of the pumping slots.

4. Photon induced desorption

4.1 Measurements at 77 K

The desorption yield measured with the beam screen at a temperature of 77 K is shown in Figure 3 as a function of the photon dose. Compared with equivalent room temperature measurements, the data for H_2 are in good agreement [6].

For this experiment the molecular desorption yield, h was calculated from the partial pressure increase making the assumption that the beam screen surface had a vanishing net pumping effect, i.e. that its surface coverage of adsorbed molecules had reached an equilibrium with the gas phase. Therefore, the only contribution to the pumping of the system was that given by the conductance of the pumping slots in the beam screen leading to adsorbtion on the 3 K surface of the cold bore. This assumption should be well satisfied for H_2 , but may not be a good approximation for the other gases.

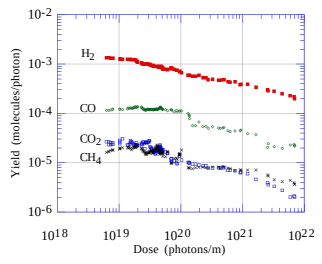


Figure 3: Desorption yields for the dominant gas species as a function of the photon dose for a beam screen temperature of 77 K.

Indeed, when comparing with room temperature results, CO and $\rm CO_2$ show a significantly lower desorption yield [7]. Whether this effect is real, i.e. the desorption of CO and $\rm CO_2$ molecules is reduced at cryogenic temperature, or can be attributed simply to a larger sticking probability on the 77 K beam screen, is uncertain. Further studies are required to confirm this observation.

4.2 Measurements at 4.2 to 10 K

As a sequel to the first measurements at 77 K, a new beam screen was installed and exposed to synchrotron radiation at a temperature which could have ranged, along its length, between the limits of 4.2 and 10 K (runs #1 and #2). During runs #3 and #4, the temperature of the beam screen was again raised to 77 K and in addition, for run #4 also the critical energy of the photon spectrum was increased to 284 eV (the value of previous measurements for the SSC) .

Throughout the experiment, individual measurements were performed by taking the differences of the pressure readings obtained with and without photon irradiation. This procedure was adopted because the pressure increases were small compared to the base pressure and because any increase of the base pressure due to a contribution from the vapour pressure of adsorbed gas molecules would have been readily detected. The data presented in Figure 4 illustrate the increase of the dynamic pressure, while the constant level of the 'photon-OFF' pressure suggests that the vapour pressure remained below the instrumental background of approximately 4×10^{-10} Torr throughout the 4 individual runs.

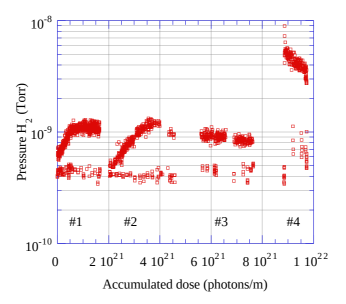


Figure 4: Summary results for the room temperature hydrogen pressure during photon exposure between 4.2 to 10 K (#1 and #2) and at 77 K (#3 and #4) showing the difference between photons On and Off. The lower points correspond to photons Off. For run #4, the critical energy of the synchrotron radiation spectrum was increased from 50 eV to 284 eV. Here the H₂ pressure increase, as measured at room temperature, has been normalised to a beam current of 300 mA in the VEPP-2M storage ring.

The molecular desorption yield calculated for H_2 is shown in Figure 5 as a function of the photon dose.

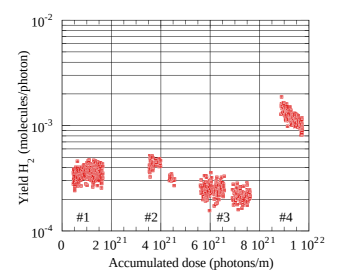


Figure 5: Summary results of the molecular desorption yield for H₂ during the 4 exposures to synchrotron radiation. (See Figure 4 for comments)

These values have been derived from the quasi steady state values in Figure 4 for which it was assumed that the screen surface had reached saturation and therefore the net pumping speed was given by the known conductance of the slots in the beam screen. Inspection of the data in Figure 5 shows a slow cleaning-up of the beam screen throughout the runs #1 to #4. It should be noted that the desorption yield at 77 K during run #3 shows no increase with respect to the 10 K measurements while the increased yield during run #4 is consistent with a linear dependence on the critical photon energy [8].

The first 2 runs #1 and #2 give a very clear illustration of the gradual increase of the dynamic pressure due to the build-up of adsorbed gas on the beam screen and the saturation level imposed by the constant pumping speed through the slots in the beam screen. Between run #1 and run #2 the beam screen was warmed-up to 77 K while maintaining the surrounding cold bore at its constant temperature of 3 K. Figure 6 shows the two successive runs.

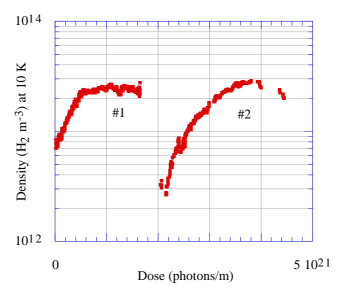


Figure 6: Evolution of the molecular density of H_2 during the photon exposure. The temperature of the beam screen is ≤ 10 K. The beam screen was warmed-up to 77 K between the two runs. After subtracting a constant background (equivalent to $4x10^{-10}$ Torr at room temperature, see Fig. 4) the H_2 density has been re-normalised to a VEPP-2M beam current of 60 mA to give a photon flux equivalent to 53 mA in LHC.

During the temporary warm-up to 77 K between run #1 and run #2 accumulated H_2 is desorbed from the screen and is permanently adsorbed on the outer 3 K surface. Therefore, run #2 is a repetition of run #1 with the only exception that the surface has been cleaned to a small degree by the additional photon exposure (e.g. see Figure 3 for the expected reduction of the molecular yield).

6. Dynamic model for H₂

The dynamic behaviour of hydrogen observed in this experiment may be described very closely by assuming a linear model for the interaction between the volume density, n(m⁻³), and the surface density, q(m⁻²), of adsorbed molecules. Apart from the replacement of ion induced desorption by photon stimulated desorption, this model is very similar to the analysis of the dynamic pressure in the ISR vacuum system [9]. The volume and the surface gas interact according to the following set of linear equations

$$V\frac{dn}{dt} = q - a \ n + b \ \theta$$

and

$$F\frac{d\theta}{dt} = c \ n - b \ \theta$$

Here, V is the volume and F is the net wall area per unit length of the vacuum system excluding the area of the pumping holes. The parameters a, b, c and q are assumed to be constant.

q represents a constant source of gas, here in particular the photon induced desorption rate determined by the product of the desorption yield, h, and of the photon flux, $\dot{\Gamma}$, hence q = h $\dot{\Gamma}$ (molecules s⁻¹ m⁻¹).

a describes the total pumping on the surface of the wall and through the area of the holes in the beam screen. It can be expressed as

$$a = \frac{1}{4} \bar{v} F(s+f)$$
 (m² s⁻¹)

where \bar{v} is the average molecular velocity, s the sticking probability of the molecules on the beam screen and f is the fraction of the total surface of the beam screen with pumping holes, f<<1.

b represent the gas source originating from the adsorbed surface phase, θ , by thermal and by photon induced desorption.

The thermal contribution [10] may be expressed as the molecular vibration frequency $n_0 = 10^{13} \text{ s}^{-1}$ multiplied with a Boltzmann factor $e^{-E_{kT}}$ containing the activation energy E.

The photon induced desorption from the adsorbed phase does not represent a source of new gas molecules but simply re-cycles molecules adsorbed on the inner side of the beam screen. It is expressed as $k\,\dot\Gamma_{,}$ where k is the re-cycling cross-section in m^2 per photon, thus

$$b = F \ v_o e^{-\frac{E}{kT}} + \kappa \dot{\Gamma}. \tag{m s-1}$$

c is the rate of adsorption of gas molecules on the surface of the beam screen only. This expression contains the sticking probability of the molecules, s,

$$c = \frac{1}{4} \overline{v} s F. \qquad (m^2 s^{-1})$$

This model contains a small number of free parameters and implies a linear adsorption isotherm, i.e. Henry's law applied to a limited range of interest

$$\theta = \frac{c}{b} n = \frac{\frac{1}{4} \overline{v} s F}{F v_{a} e^{-E/kT} + \kappa \dot{\Gamma}} n.$$

Here θ (n) depends on the re-cycling cross-section as well as on the photon flux. Therefore, the isotherm and the equilibrium surface coverage for a given volume gas density will depend on the specific conditions (i.e. on the beam current) during the experiment which effectively reduce the sojourn time of the physisorbed molecules.

As is shown for run #2 in Figure 7, the results can indeed be described in a fairly satisfactory manner by this simple model by adapting the free parameters summarised in Table 1.

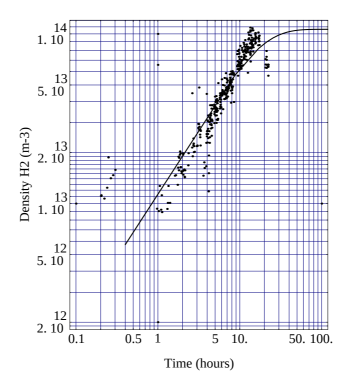


Figure 7: Comparison of the H₂ data from run #2 with the dynamic model of desorption and re-cycling of H₂ molecules. The specific parameters which have been used are summarised in Table 1.

Table 1

desorption yield	η	5x10 ⁻⁴ molecules/photon
screen temperature	T	10 K (maximum)
linear photon flux	Γ̈́	3.14×10^{16} photons/s/m
(200 mA)		-
sticking probability	S	0.6
fraction of slots	f	1.28%
monolayer capacity		3x10 ¹⁹ molecules/m²
re-cycling	κ	$5 \times 10^{-21} \mathrm{m}^2$
cross-section		
activation energy	E	800 cal/mole
		0.035 eV/molecule
specific surface area	F	0.14 m
specific volume	V	1.3x10 ⁻³ m ²

Parameters shown in bold face have been adapted to obtain the curves in Figures 6-8. The fraction of slots has been corrected for the Clausing factor due to the thickness of the beam screen.

The model provides in addition to the volume density also the surface density of gas, q, shown in Figure 8. There it can be seen that the final coverage obtained remains well below a monolayer.

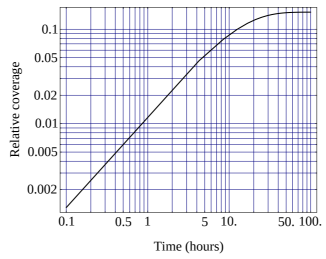


Figure 8: Surface density of H₂ molecules normalised to a monolayer coverage as a function of time.

As a further result, Figure 9 gives the calculated volume densities as a function of time with photons On and photons Off. The lower curve (photons Off) corresponds to the equivalent measurements in Figure 4 and gives the vapour pressure of the adsorbed gas. Considering that the lowest detectable pressure rise in the system corresponds to a density of approximately $8x10^{12}$ molecules m⁻³, it is not surprising that the calculated increase in vapour pressure could not be observed during the runs.

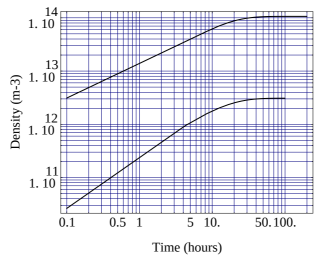


Figure 9: Upper curve shows the gas density with photon exposure and lower curve shows the gas density without photons.

7. LHC performance

The test system and the beam screen used for these measurements very closely reproduce the conditions in LHC. At 230 mA in VEPP-2M the linear photon flux in the test system is only about a factor of 2 lower than that in the LHC at 7 TeV and with the nominal beam current of 536 mA. To obtain an estimate of the vacuum performance during LHC operation, it was therefore sufficient to scale the linear

photon flux since running time in hours and H₂ density in the beam screen depend both linearly on the photon flux. The scaling of runs #1 and #2 to LHC conditions with an initial beam current of 53 mA but without modifying the arrangement of pumping slots is shown in Figure 10. Here the dose scale has been converted to running time in hours to illustrate the short time necessary to reach the stationary density due to the density limiting effect of the pumping holes. Without holes, the density would continue to rise and would exceed the maximum tolerable level within a few days of operation.

The lifetime limit of 10^{15} molecules m⁻³ for H₂ is determined for LHC by the tolerable rate of particle loss from the beam and corresponds to a beam lifetime of approximately 100 hours. So far only H₂ has been taken into account. In a real vacuum system, the contribution of all other gas species has to be added in proportion of their respective cross sections and thus the apparently comfortable vacuum performance shown in Figure 9 may not be achieved. For this reason and in order to meet the vacuum requirements with the 10 times larger nominal beam current (530 mA), the LHC beam screen will need a substantial amount of in-situ cleaning by synchrotron radiation in the same way as for electron machines.

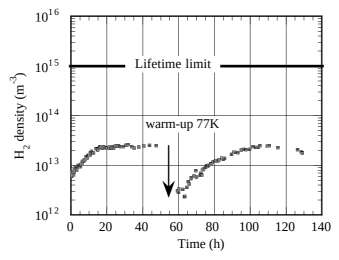


Figure 10: Dynamic H_2 pressure in LHC with an initial beam current of 53 mA derived from the results of the first 2 runs.

Important for the design of the LHC vacuum system will be the total amount of H_2 desorbed. This quantity may be estimated from an extrapolation of the H_2 data in Figure 3 where it can be seen that the yield as a function of photon dose may be approximated by

$$\eta = \eta_o \left(\frac{D}{D_o}\right)^{-\frac{1}{3}}$$

with $\eta_o = 10^3$ molecules/photon and $D_o = 2 \times 10^{19}$ photons m⁻¹. The accumulated photon dose in LHC during the first year will be of the order of 10^{23} photons m⁻¹ and thus one can estimate that the equivalent of 2.6 monolayers of H₂ will be desorbed from the beam screen. During the second year, at the full nominal beam current, the equivalent of 10.5 monolayers will be desorbed. This total quantity of gas should not pose a problem within the present design of the LHC

vacuum system since the molecules can diffuse through the pumping holes and be adsorbed on the 1.9 K cold bore with a negligible vapour pressure.

Acknowledgements

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