



EFDA-JET-CP(03)04-03

C. H. Skinner, J. P. Coad and G. Federici and JET EFDA Contributors

Tritium Removal from Carbon Plasma Facing Components

Tritium Removal from Carbon Plasma Facing Components

C.H. Skinner¹, J.P. Coad² and G. Federici³ and JET EFDA Contributors*

 ¹Princeton Plasma Physics Laboratory, Princeton, NJ USA
²EURATOM/UKAEA Fusion Association, Culham Science Centre, Abingdon, OX14 3DB, UK
³ITER Garching Joint Work Site, Garching, Germany
* See annex of J. Pamela et al, "Overview of Recent JET Results and Future Perspectives", Fusion Energy 2000 (Proc. 18th Int. Conf. Sorrento, 2000), IAEA, Vienna (2001).

> Preprint of Paper to be submitted for publication in Proceedings of the 10th International Workshop on Carbon Materials for Fusion Application (Jülich, Germany, 17-19 September 2003)

"This document is intended for publication in the open literature. It is made available on the understanding that it may not be further circulated and extracts or references may not be published prior to publication of the original when applicable, or without the consent of the Publications Officer, EFDA, Culham Science Centre, Abingdon, Oxon, OX14 3DB, UK."

"Enquiries about Copyright and reproduction should be addressed to the Publications Officer, EFDA, Culham Science Centre, Abingdon, Oxon, OX14 3DB, UK."

ABSTRACT.

Tritium removal is a major unsolved development task for next step devices with carbon plasma facing components. The 2-3 order of magnitude increase in duty cycle and associated tritium accumulation rate in a next step tokamak will place unprecedented demands on tritium removal technology. The associated technical risk can be mitigated only if suitable removal techniques are demonstrated on tokamaks before the construction of a next step device. This article reviews the history of codeposition, the tritium experience of TFTR and JET and the potential impact of tritium removal on ITER's planned physics schedule. The merits and shortcomings of various tritium removal techniques are discussed with particular emphasis on oxidation and laser surface heating.

1. INTRODUCTION

Fast and efficient tritium removal will be essential in any next step magnetic fusion reactor with carbon Plasma Facing Components (PFCs). At the same time the necessary technology still needs to be established on a working tokamak, in conspicuous contrast to almost every other enabling technology, such as remote handling and superconducting magnets. This is a major unsolved development task for next step devices with carbon PFCs. When operating with tritium fuel both the Tokamak Fusion Test Reactor (TFTR) and the Joint European Torus (JET) were able to remove sufficient tritium to stay within inventory limits, however in a next step device the orders-ofmagnitude increase in duty cycle will greatly increase the rate of tritium accumulation. Modelling for the International Thermonuclear Experimental Reactor (ITER) with carbon PFCs predicts that the in-vessel tritium inventory limit could be reached after a few weeks of DT plasma operations and tritium removal at unprecedented speed and efficiency will be required to maintain a credible plasma physics program. The magnetic and radiological environment and relative inaccessibility of parts of the divertor place severe demands on the removal technology. Simply predicting the amount and location of codeposited tritium in existing divertor machines is a challenge for state-ofthe-art models. Of course, the use of tungsten PFCs would avoid this issue and tungsten is planned as an alternative for ITER if appropriate tritium removal technology cannot be demonstrated. However tungsten carries its own risks. Maintaining plasma purity with high-Z materials is a concern, and depending on the frequency and severity of ELMs and disruptions, erosion due to tungsten melt layer loss could prematurely terminate the useful lifetime of the divertor and also disrupt the planned physics program.

Extensive references on this area are available. A comprehensive review of plasma material interactions and their implications for next step tokamaks including tritium removal techniques was presented in ref. [1]. Tritium retention and removal in the two tokamaks to use tritium fuel; TFTR [2-10] and JET[11-14] has been extensively reported. Hydrogen isotope retention and recycling data is reviewed in [15]. Thermal oxidative techniques for removing hydrogen isotopes were reviewed in a paper at the 9th Carbon workshop[16]. Detection and removal of carbon debris is described in ref. [17]. Tritium issues in ITER are reviewed in [18-22]. A survey of techniques to detritiate tiles after their removal from fusion machines is in refs. [23,24]. This paper will begin with a short

history of hydrogen-isotope / carbon interactions in a plasma, from the successful use of graphite limiters on PLT to the experience with tritium retention in JET and TFTR and then focus on tritium removal by oxidation and by laser surface heating.

2. CO-DEPOSITION

Codeposition of hydrogen isotopes with carbon is the tangible result of the interaction of plasmas with carbon PFCs (figure 1). Carbon has a low Z and excellent mechanical properties at high temperature, ideal parameters for next step DT burning reactors, however carbon PFCs will not be acceptable without proven methods to rapidly and efficiently remove the codeposited tritium. Codeposition was not an issue in the early days of fusion research when stellerators used metal limiters. Plasma performance advanced with the development of tokamaks, however initial attempts to reach high temperature plasmas in the Princeton Large Torus (PLT) were stymied by the increase in core radiation from tungsten impurities[25]. Changing the limiter material from tungsten to carbon in 1978 enabled the successful attainment of low collisionality plasmas at ion temperatures of 5.5keV and carbon has been the favored plasma facing material ever since.

The advantages of a carbon plasma facing surface led to the development of carbonization techniques where a carbon layer was deposited on metals by a radio frequency assisted DC-glow discharge technique[26]. A reduction in tritium inventory with carbonized walls as compared to stainless steel was anticipated due to lower hydrogen isotope mobility. However, it became clear from analysis of graphite limiter tiles after the operation of the JET[27] and TFTR[28] tokamaks with carbon PFCs, that codeposition trapped hydrogen isotopes much more effectively than implantation and the amount retained could increase indefinitely without saturation. Widespread recognition of codeposition and its consequences was slow in coming. A 1984 data compendium discussed ion implantation, diffusion, reflection and desorption, but made no mention of codeposition[29]. A 1997 edition of a textbook on tokamaks [30] strangely omits codeposition from a discussion of tritium behavior in tokamaks. The ITER Physics Basis [31] mentions the need for tritium removal technology, but does not discuss removal methods even at a conceptual level, nor the potentially serious impact on the planned physics program of the interruptions required for tritium removal.

3. TRITIUM RETENTION IN TFTR, JET AND ITER.

Fusion research attained a major milestone with the use of tritium fuel on TFTR and JET, producing 10 and 16MW of fusion power respectively. In TFTR, 5g of tritium were injected into circular plasmas over a 3.5 year period, mostly by neutral beam injection [32]. In JET, 35g were injected into diverted plasmas over a 6 month campaign, mostly by gas puffing [33]. In TFTR the bumper limiter provided a large source of eroded carbon and an average of 5% of tritium was co-deposited on the limiter and vessel wall during plasma operations. Only a small area of the codeposit flaked off [34]. In JET the wall is a net erosion area, and co-deposition occurs principally in shadowed parts of the inner divertor, with heavy flaking. The retention rate during the DTE1 campaign was

40%. In both machines, tritium retention was initially high following a change from deuterium to tritium gas puffing, due to isotope exchange with deuterium on plasma facing surfaces (dynamic inventory). The contribution of codeposition is lower but cumulative, and is revealed by including periods of D fuelling that reversed the T/D isotope exchange.

The tritium retention rate varied depending on the circumstances (fueling method, history of plasma discharges, tritium inventory in the vessel etc...). For example from the beginning of DT operations in December 1993 until August 1995, 1.9g of tritium entered the TFTR torus and approximately 40% or 0.7g was retained. This fraction was similar to the earlier deuterium measurements. In September 1995 L-mode discharges were fueled by gas puffing in contrast to the prior neutral beam fuelled supershots. Tritium was taken up on the previously depleted walls such that of the 1.1g T supplied for these experiments only 0.09g was pumped out, a retention fraction above 90%. After a few days the in-vessel inventory reached 1.8g T, close to the 2.1g administrative limit and tritium removal was necessary for continued plasma operations. The internal configuration of TFTR was not changed in this period and the overall average TFTR tritium retention rate during plasma operations (51%) was in line to the prior experience with deuterium [35] and consistent with first principles calculations[9,32]. However one previously suggested method to remove tritium by helium ohmic discharge conditioning[35] was not useful in practise.

The percentage of deuterium fuel retained in JET increased with increasing coverage of graphite wall tiles [36]. On an 'all carbon' JET, gas balance measurements show that deuterium was retained at a rate of 40% of the input, principally by codeposition [37]. Some of this was lost during venting etc and PFC analysis showed long term retention of deuterium by codeposition at a rate of 17% of the deuterium input[38]. The introduction of beryllium PFCs increased the deuterium required to fuel the plasma by about a factor of four, but the absolute amount of retained deuterium did not increase significantly. Tritium behaviour in JET DTE1 experiments was surprisingly different to that in the earlier PTE experiments. The tritium inventory increased a factor-of-two faster than expected[12] peaking at 11.5g T with more than half of the tritium on site trapped in the vessel. This increase is attributed to changes in the temperature and geometry of the divertor however the absolute amount of tritium retained is still not quantitatively understood[39]. Pulsing in deuterium reduced the inventory to 6.2g of tritium but was ineffective in reducing it further. From the PTE experience 1g of tritium was expected to be retained on tiles, but measurements showed that the DTE1 levels on the tiles were lower. In contrast, the tritium outgassing rate was an order of magnitude higher than expected. The major difference in DTE1 was the formation of films with high (~ 0.8) D/ C ratio on the divertor louvers and subsequent flaking and accumulation in the sub-divertor[13]. This relatively inaccessible location hampered efforts at tritium removal. An additional concern is the discovery of tritium trapped in the bulk of CFC tiles where it will be extremely difficult to remove[40].

Deuterium-tritium operations are scheduled for ITER after 3 years of hydrogen, and 1 year of deuterium plasmas. ITER will be fuelled by approximately 120g T per 400s pulse and of that, 2-5g T is anticipated to be trapped in the vessel principally by codeposition with eroded carbon[19]. The in-vessel tritium inventory is limited to 350g T to avoid evacuation at the site boundary in the case

of the worst credible accident. Independent of safety considerations, tritium is expensive and the supply is limited so it is important to avoid inventory build up in inaccesible locations. In the worst case, the ITER in-vessel inventory limit could be reached after just one week of DT operations (Fig.2). This estimate is uncertain due to lack of code validation in detached plasma regimes (in fact the same code underestimated tritium retention in JET by a factor ×40 [39]). Retention could be higher if the ITER outer wall is a carbon deposition area [41] or if there is significant carbon erosion by type 1 ELMS [42]. On the other hand retention could be lower if beryllium impurities impede chemical erosion of carbon near the divertor strike points and hence tritium codeposition. Formation of mixed materials could make tritium removal more challenging. Once the tritium inventory limit is approached, DT plasma operations will be terminated and not allowed to restart until substantial amounts of tritium are removed from the vessel. The delay this entails is highly uncertain since no relevant method to remove tritium has been established on a working tokamak. Despite these uncertainties it is very clear that fast and efficient tritium removal will be absolutely necessary for ITER with carbon PFCs to meet its physics mission.

4. TRITIUM REMOVAL

The scale-up in the ITER duty cycle places huge demands on any detritiation technique. ITER operations envisages 2 shifts (14 hours/day) 5 days/week, 3 weeks/month, 8 months / year with 70% availability to reach the design goal of approximately 2,000 pulses/year [43]. Each pulse is 400s long and occurs every 33 minutes. The current projected tritium codeposition rate is 2-5g T per pulse, principally from chemical erosion of a 20m² area in the divertor and would leave 50 - 127g T to be removed every overnight shift! Techniques such as glow discharge cleaning that are not compatible with the 5T toroidal field will not be suitable. The peak codeposition rate for the ITER outer divertor is 5nm/s [44] would result in a peak codeposited layer thickness growth of 50microns / day or 0.25mm / week. To remove the tritiated codeposit in a routine tritium cleanup operation taking half the overnight shift a removal rate of 10 microns/hr is required. The ITER situation is contrasted with prior experience on TFTR and JET in Table 1.

Several methods to remove tritium were investigated during the ITER engineering design activity and are reviewed in [1,18,19]. The methods may be grouped in two classes. One option is to break the a-C:T chemical bond by heating to high temperatures, or by UV or chemical means such as isotope exchange. The tritium is then desorbed as T2 or DT gas and pumped out. The other option is to remove the whole codeposit by oxidation or by ablation with a pulsed laser or flashlamp. Potential techniques must be compatible with the 5 Tessla field as the time required to cycle the field off and on is prohibitive. It should also be compatible with the gamma field from activated components. After 20 years of operation this is estimated to be 10,000Gy/h at 10⁴s after shutdown [45]. Tokamak demonstrations are important to demonstrate that the released tritium is recoverable. For example the technique should not produce reactive radicals that could be reabsorbed before exiting the torus. Debris produced by ablative methods will need to be efficiently collected [46].

The merits and shortcomings of several techniques are listed in Table 2. In this paper we focus on two techniques – oxidation and transient heating by a scanning laser.

4.1 HYDROGEN ISOTOPE REMOVAL VIA OXIDATION

Hydrogen isotopes are released from carbon at elevated temperatures. Early laboratory experiments showed deuterium was released from both codeposited and ion implanted carbon samples on heating to 800°C in vacuum but this temperature is much higher than typical PFCs. Fortunately in air the release temperatures are much lower: $\approx 300 - 400^{\circ}$ C [47]. Comparison of the effect of H₂, O₂, and H₂O showed oxygen was the most effective gas in removing deuterium [48] and the reaction mechanism was simple thermal oxidation leading to emission of D₂O, CO₂ and CO [49]. The removal rate varied with the film structure, codeposited films from ASDEX-Upgrade being removed much faster than films grown in an RF glow-discharge[50]. Tests on codeposited films from TEXTOR showed oxygen (300-600 Pa) at 250 – 400°C was effective[51]. Soft films were oxidised at lower temperatures than hard films[52]. The removal rate of codeposited films on tiles removed from TFTR, JET, DIII-D and ASDEX-U ranged from 0.1 – 50µm/h on exposure to oxygen at 250 and 350°C [53, 54] with faster erosion rates for thicker deposits (see Fig.3). The structure of codeposits is more open and porous than the underlying graphite. The erosion rate for manufactured graphite is conveniently two to three orders of magnitude slower so that while the codeposit is removed and tritium released, the thickness of the graphite tile is hardly affected.

Oxidative removal of tritium was very helpful in enabling TFTR and JET to control the tritium inventory. Introducing air into the room temperature TFTR torus followed by pumping released 0.2 g T in three cycles. This operation took 2 days for a fill at 0.99 bar, the removal rate being limited by the time required to process the tritium. At the end of plasma operations 0.2 - 0.8 bar of air was introduced with the torus at 150° C and 0.14 g T released. On JET a similar exercise removed 0.6 g of tritium (about 10% of the inventory).

An important test of the efficacy of oxygen to remove hot codeposited layers and release hydrogen isotopes was performed on TEXTOR [55]. Oxygen was introduced into the torus at pressures between 0.007 and 0.3mbar and wall temperatures between 227°C and 427°C. Part of the oxygen reacted to form compounds on the wall, part reacted to produce CO and CO₂. gases which were pumped out. Hydrogen isotopes incorporated in codeposits were released as water molecules (not as hydrogen molecules or hydrocarbons) but some of the released water was reabsorbed on the vessel walls. After oxygen exposure, He and D glow discharges were needed to restore wall conditions suitable for plasma operations and boronization was performed to reduce the oxygen impurities to the normal low levels. A quantitive estimate of the efficacy of hot oxygen to enable the pump out of hydrogen isotopes was not possible and needs more experiments and better understanding of water absorption and desorption behavior from the entire wall.

This technique can access all areas including gaps, shadowed regions and areas not in line of sight to the plasma. However there are serious obstacles. The temperature of the ITER PFCs is limited to 240° C (the maximum temperature of the pressurized water coolant) and oxidation at this

temperature is an order of magnitude slower [53, 56]. Oxygen is known to decondition the plasma facing surfaces and a long clean- up time will be required to remove the oxygen before good plasma performance is restored. The oxygen may damage in-vessel components. The potential for mixed material effects in ITER adds further uncertainty. Tungsten[57] or boron [58] impurities were found to inhibit the oxidation of hydrocarbon films possibly through masking of the film surface. The resulting tritium oxide is 27,000 times more hazardous than elemental tritium and processing the exhaust requires a large investment in the tritium plant. For this technique to be a viable candidate more tokamak demonstrations with oxygen at the ITER conditions are clearly required, however this proposal is not looked on with favor by tokamak operators because of the known deleterious effects of oxygen on plasma performance.

4.2 HYDROGEN ISOTOPE REMOVAL BY LASER SURFACE HEATING

Recently a novel method for removing tritium was demonstrated. The operating principle is robust - it is well known that tritium is thermally desorbed at 700°C temperatures, but heating the whole tokamak to that temperature is impractical. However most of the tritium is trapped in codeposit layers and that can be heated by a rapidly scanning laser beam in vacuum or an inert atmosphere without the deleterious effects of oxidation [59]. Tabletop experiments with a scanning Nd laser have since demonstrated efficient removal of tritium from JET and TFTR tile samples by this method [60, 61]. A 325 W continuous wave Nd laser beam was steered by two orthogonal mirrors and focused inside a chamber containing the tile sample. A serpentine raster pattern with line spacing 0.5mm was used to cover the tile surface at a scan speed of 1m/s covering an area of 5cm²/s. The duration of heating pulse was typically 1.6 ms and incident energy up to 0.2MJ/m². The surface temperature rise was much higher for codeposits than the bare tile material, because the codeposit was less thermally conductive or not in good thermal contact with the substrate[62]. The transiently high surface temperatures (over 2300°C in some cases) readily released the tritium which was measured by an ion chamber in a closed loop circulation system. Tritium remaining after the laser scan was released by baking the codeposit in air at 500°C for 15 mins and the efficiency of the laser detritiation derived. Fig. 4 shows the results from seven samples scanned in a variety of conditions, not all optimal. The detritiation efficiency is high, up to 87%, and is the highest on the heaviest codeposits.

The atmosphere of the chamber is circulated continuously and the tritium concentration did not change after the scan indicating that reabsorption back on to the tile or isotope exchange with water on the interior chamber surface was minimal. The codeposit remained after the temperature excursion, though the surface changed color from brownish to dark gray.

For tokamak applications the laser would remain outside the vessel and be coupled via a fiber optic to a scan head inside. Fiber optic coupling was demonstrated in the above experiments and design considerations for the application to ITER are discussed in ref.[63]. Industrial fiber optic coupled Nd lasers are available at powers up to 6kW and the energy required to heat a 200 micron surface of 50m² area to 2,000°C could be delivered over 3 hours. Activation will generate a strong

gamma field inside ITER, however recently developed radiation resistant optical fibers have sufficient transmission. The scanner will be constructed from non-magnetic materials to be compatible with the 5 Tessla field. Novel scan head designs have been proposed to allow coverage of hard-to-access areas between tiles and in hidden areas. Development of prototypes and demonstrations of in-situ detritiation of a tokamak are needed but are currently unfunded.

CONCLUSIONS.

ITER's ability to address burning plasma physics depends on the successful resolution of serious plasma material interaction issues. The change in ITER pulse length and duty cycle is larger than the change in any plasma parameter and poses severe challenges in tritium management and material erosion. Just as suitable plasma physics regimes are thoroughly explored on other tokamaks before being planned for ITER, all the candidate materials for ITER should be fully tested in a large tokamak and any issues addressed with dedicated R&D before ITER PFC procurement decisions are made. Without this step, ITER will inevitably begin operation as very expensive test bed for plasma material interactions. Much progress has been made in traditional plasma physics and fusion technology fields, however 'housekeeping' areas such as tritium removal have not fared so well even though they are essential to any fusion reactor with carbon PFCs. Arguably the biggest technical risks in a burning plasma experiment are related to the choice of PFCs. Public attitudes toward tritium and the prospective \approx \$100,000 per hour cost of any unforeseen interruption in the ITER physics program are likely to make the lack of established tritium removal technology an increasingly sensitive point in the public support of fusion research. A scale up of R&D effort on these issues is urgently needed to establish high-confidence solutions on tokamaks so that fusion can meet its promise as an attractive environmentally acceptable energy source for mankind

ACKNOWLEDGEMENTS

The authors gratefully acknowledge stimulating discussions with our colleagues in this field, This work is partially funded by U.S. DOE Contract Nos. DE-AC02-76CH0307, by the UK Department of Trade and Industry, and as an account of work undertaken within the framework of ITER Transitional Arrangements (ITA). These are conducted by the Participants: Canada, China, the European Atomic Energy Community, Japan, the Russian Federation, and the USA under the auspices of the International Atomic Energy Agency. The views and opinions expressed herein do not necessarily reflect those of the Participants to the ITA, the IAEA or any agency thereof. Dissemination of the information in this paper is governed by the applicable terms of the former ITER-EDA Agreement.

REFERENCES

- G. Federici, C. H. Skinner, J. N. Brooks, J. P. Coad, C. Grisolia, A. A. Haasz, A. Hassanein, V. Philipps, C. S. Pitcher, J. Roth, W. R. Wampler, D. G. Whyte, Nucl. Fus., 41 (2001) 1967.
- [2]. C. H. Skinner et al., J. Vac. Sci. Technol. A14 (1996) 3267.

- [3]. D. Mueller et al., Fus. Tech., **30** (1996) 840.
- [4]. D. Mueller et al., J. Nucl., Mater., **241-243** (1997) 897.
- [5]. D. Mueller et al., Proceedings of the 17th IEEE/NPSS Symposium on Fusion Engineering, San Diego, October 6-10, 1997, p279, IEEE, Piscataway, NJ, USA (1998).
- [6]. A Nagy et al., Proceedings of the 17th IEEE/NPSS Symposium on Fusion Engineering, San Diego, October 6-10, 1997, p**317**, IEEE, Piscataway, NJ, USA (1998)
- [7]. C. H. Skinner et al., Proceedings of the 20th Symposium on Fusion Technology, Marseille, Sept. 7-11, 1998 v.1, p153-156, Association EURATOM-CEA, Cadarache, France, September 1998.
- [8]. C. H. Skinner et al., J. Nucl. Mater., 226-269 (1999) 940.
- [9]. C. H. Skinner et al., J. Nucl. Mater., 290-293 (2001) 486.
- [10]. C. H. Skinner et al., 28th EPS Conference on Controlled Fusion and Plasma Physics Madeira, Portugal, 18-22nd June, 2001 Europhysics Conference Abstracts, 25A (2001) 1621-1624
- [11]. P. Andrew et al., J. Nucl. Mater., 266-269 (1999) 153.
- [12]. P. Andrew et al., Fus. Engin. & Des., 47 (1999) 233.
- [13]. A. T. Peacock et al., Fus. Eng. & Des., 49-50 (2000) 745.
- [14]. J. P. Coad et al., J. Nucl. Mater., **290-293** (2001) 224.
- [15]. R. A. Causey, J. Nucl. Mater., **300** (2002) 91.
- [16]. J. W. Davis and A. A. Haasz, Phys. Scripta T91 (2001) 33.
- [17]. G. F. Counsell and C. H. Wu., Phys. Scripta T91 (2002) 70.
- [18]. G. Federici et al., Fus. Eng. & Des., **39-40** (1998) 445.
- [19]. G. Federici et al., J. Nucl. Mater., 266-269 (1999) 14.
- [20]. G. Federici, J. N. Brooks, M. Iseli and C. H. Wu, Phys. Scripta T91 (2001) 76.
- [21]. G. Federici and C. H. Skinner, "Tritium Inventory in the materials of the ITER plasma-facing components" Springer Verlag, Berlin-Heidelberg 2002.
- [22]. G. Federici et al., J. Nucl. Mater., 313-316 (2003) 11.
- [23]. N Bekris, C. Caldwell-Nichols, L Doerr, M. Glugla, R.-D. Penzhorn, H. Zeigler., J. Nucl. Mater., 307-311 (2002) 1649.
- [24]. A. C. Bell, J. Williams, J. D. Nielson, and A. Perevezentsev, Fus. Sci., Technol., 41, (2002) 626.
- [25]. H. Eubank et al., in Plasma Physics and Controlled Nuclear Fusion Research 1978 (Proc. 7th Int. Conf. Innsbruck, 1978) Vol. 1 IAEA Vienna (1978), p. 167.
- [26]. J. Winter. J. Nucl. Mater., 145-147 (1989) 131.
- [27]. H. Bergsaker et al., J. Nucl. Mater., 145-147 (1987) 727.
- [28]. W. R. Wampler et al., J. Vac. Sci. Technol. A6 (1988) 2111.
- [29]. R. L. Langley et al., Data Compendium for Plasma-Surface Interactions Nuclear Fusion Special Issue (1984). IAEA, Vienna 1984 p.28.
- [30]. J. Wesson "Tokamaks" Clarendon Press-Oxford 1997 p.463.
- [31]. Nucl. Fus., **39** (1999) 2137.
- [32]. C. H. Skinner et al., J. Nucl. Mater., 266-269 (1999) 940.

- [33]. P. A. Andrew et al., Fusion Engineering Design 47 (1999) 233.
- [34]. C. H. Skinner et al., Nucl., Fus., **39** (1999) 1081.
- [35]. A. E. Pontau et al., Fus. Engin & Des., 10 (1989) 365.
- [36]. J. P. Coad et al., J. Nucl. Mater., 162-164 (1989) 533.
- [37]. R. Sartori, G. Saibene et al, J. Nucl. Mater., 176&177 (1990) 624.
- [38]. J. P. Coad., J. Nucl. Mater., 226 (1995) 156.
- [39]. J. N. Brooks, A. Kirchner, D. G. Whyte, D. N. Ruzic and D. A. Altman, J. Nucl. Mater., 313-316, (2003) 424.
- [40]. R.-D. Penzhorn, N. Bekris, U. Berndt, J. P. Coad, H. Zeigler and W. Naegele, J. Nucl. Mater., 288 (2001) 170.
- [41]. A. S. Kukushkin, H. D. Pacher, D. Coster, G. W. Pacher, D. Reiter, 30th EPS Conference on Controlled Fusion and Plasma Physics, St. Petersburg, Russia July 7- 11th 2003, Europhysics Conference Abstracts
- [42]. G. Federici, A. Loarte, G. Strohmayer, "Assessment of erosion of the ITER divertor targets during type I ELMs", to appear in Plasma Phys. Control. Fusion (2003).
- [43]. E3 Physics Operations Working Group Final Report 2002 Fusion Energy Sciences Summer Study (Snowmass 2002, unpublished).
- [44]. J Brooks, personal communication.
- [45]. H. Iida, personal communication.
- [46]. Ph. Delaporte et al., Appl. Surface Science 197-198 (2002) 826.
- [47]. R. A. Causey, W. R. Wampler and D. Walsh, J. Nucl. Mater., 176-177 (1990) 987.
- [48]. S. Chiu and A. A. Haasz, J. Vac. Sci. Technol., A9 (1991) 747.
- [49]. A.A. Haasz, S. Chiu, J. E. Pierre, and Y. I. Gudimenko, J. Vac. Sci. Technol., A14 (1996) 184.
- [50]. W. Wang, W. Jacob and J. Roth, J. Nucl. Mater., 254 (19997) 66.
- [51]. S. Alberici et al., J. Nucl. Mater., 258-263 (1998) 764.
- [52]. K. Maruyama, W. Jacob, J. Roth., J. Nucl. Mater., 264 (1999) 56.
- [53]. A. A. Haasz and J. W. Davis., J. Nucl. Mater., 256 (1998) 65.
- [54]. J. W. Davis and A. A. Haasz, J. Nucl. Mater., 266-269 (1999) 478.
- [55]. V. Philipps et al., J. Nucl. Mater., 266-269 (1999) 386.
- [56]. W.M. Shu, C.A. Gentile, C.H. Skinner, S. Langish, M.F. Nishi, Fus. Engin. & Des., 61-2 (2002) 599.
- [57]. J. W. Davis, C. G. Hamilton, A. A. Haasz, R. G. Macaulay-Newcombe, J. Nucl. Mater., 305 (2002) 66.
- [58]. J. W. Davis, P. B. Wright, R. G. Macaulay-Newcombe, A. A. Haasz, C. G. Hamilton, J. Nucl. Mater., 290-293 (2001) 66.
- [59]. C. H. Skinner, H Kugel, D Mueller, B. L. Doyle, and W. R. Wampler, Proceedings of the 17th IEEE/NPSS Symposium on Fusion Engineering, San Diego, October 6-10, 1997, vol. 1, pp.321-324, IEEE, Piscataway, NJ, USA (1998).

- [60]. C. H. Skinner, C. A. Gentile, A. Carpe, G. Guttadora, S. Langish, K. M. Young, W. M. Shu, H. Nakamura, J. Nucl. Mater., 301 (2002) 98.
- [61]. C. H. Skinner, N. Bekris, J. P. Coad, C. A. Gentile, M. Glugla, J. Nucl. Mater., 313-316 (2003) 496.
- [62]. C. H. Skinner, N. Bekris, J. P. Coad, C. A. Gentile, A. Hassanein, R. Resiwig and S. Willms Phys. Scripta T 103 (2003) 34.
- [63]. C. H. Skinner, C. A. Gentile, G. Guttadora, A. Carpe. S. Langish, K. M. Young, M. Nishi, W. Shu, Fusion Science and Technology 41 (2002) 716

Parameters:	TFTR experience	JET experience	ITER projections
Tritium in-vessel inventory limit	2g	20g site inventory	350g
Period of DT operations before inventory limit first approached.	Dec'93 to Sept'95 or 22 months	3 months left 11.5g in vessel	≈ 1 week (w/uncertainties)
Typical pulse duration	≤ 8s	30s	400s
Cumulative DT discharge duration before inventory limit first approached.	708 pulses 33 min	500 pulses 250 min	≈70–170 pulses 466 – 1,133 min
Time devoted to tritium removal	1.5 months*	3 months	est. ≈5h overnight or ≈14h weekend available
Fraction of tritium removed	50%	50% (prior to venting)	?

*some of this time was devoted to other maintenance activities.

JG03.712-T1

Table 1: Comparison or tritium parameters of TFTR, JET & ITER.

Technique	Merits	Shortcomings
Air/O ₂ exposure with hot walls	Technical simplicity, access to all areas	Removal rate insufficient at ITER design temperature (240 ;C). Wall reconditioning required to remove residual oxygen and water to recover plasma operation. Inhibited by tunsten or boron impurities. Potential for collateral damage. DTO expensive to process.
Plasma assisted oxidation via ECR or ICR with oxygen	Does not require vent, limited tokamak experience.	Removal rate insufficient especially in areas not line of sight to plasma Wall reconditioning required to remove residual oxygen and water to recover plasma operation. Potential for collateral damage. DTO expensive to process.
Isotope exchange with D plasmas	JET	Removal rate inadequate for ITER
Divertor to include Cold Catchers periodically heated to recover tritium		No tokamak experience. Depends on accurate predictions of location of tritium.
Photo ablation	Industrial experience	Appears incompatible with 5 Tessla field. Collection of any tritiated debris required.
Laser ablation	Laboratory experience	Technically difficult to apply inside ITER vessel. Collection of any tritiated debris required.
Thermal desorption by scanning laser	Demonstrated on JET & TFTR samples in laboratory. Fast, scalable to ITER vessel. No oxygen involved.	Tokamak demonstration needed.

JG03.712-T1

Table 2: Techniques for tritium removal





Figure 1 Microscope image of thick codeposit on TFTR graphite tile. The image covers 282 microns in the horizontal direction (reproduced with permission from Ref. 62).

Figure 2 Tritium retention in ITER, showing modeling predictions and the JET DTE1 equivalent rate. The inventory limit (shown by double line) is predicted to be reached in approximately 100 pulses.





Figure 3: Film thickness as a function of O_2 exposure time at 523, 573 and 623K for a TFTR codeposited tile specimens. Reproduced with permission from ref.[53].

Figure 4: Tritium released by laser scanning and the subsequent bake in air for samples from JET and TFTR. A variety of scan conditions were used, some not yet optimal. Non the less the results illustrate that the detritiation efficiency is high at the locations with the highest concentrations of tritium. Reproduced with permission from ref [61].