

Superiority of Electropolishing over Chemical Polishing on High Gradients

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Abstract

Electropolishing (EP) offers a smoother surface, and that makes us expect high gradients in superconducting niobium cavities. EP has a little bit complicated procedure and needs annealing to avoid the hydrogen Q-disease. On the other hand, chemical polishing (CP) is easier and cheaper, so widely used in many laboratories. For the conventional field gradients (≈ 5 MV/m) in storage rings (TRISTAN-MR, HERA), no difference was seen on the cavity performance between EP and CP. In the high gradients research for TESLA, we have to establish the surface treatment method which has a high reliability on the high gradients over than 25 MV/m and is cheaper in the preparation cost. So far, CP has offered a nice cavity performance, but a concern on "European Headache" was developed recently on the high gradients at Saclay. KEK has intensively investigated the difference between the effect of CP and EP on high gradient performance since the last SRF workshop at Saclay. In this investigation "European headache" was also seen in chemically polished cavities at KEK, in addition these cavities often showed a new Q-degradation without x-ray signals which seriously limited the high gradient. EP offers the high gradient over than 30 MV/m reliably. At the moment, EP seems to have a superiority over CP on the high gradient over than 25 MV/m.

1. Introduction

KEK has adopted electropolishing (EP) for the preparation of niobium superconducting rf cavities, while chemical polishing (CP) is widely used in the other laboratories because it is easier than EP. EP can offer a smoother surface than CP, thus makes us expect to obtain a better cavity performance. Really, KEK established the 10 MV/m level in the accelerating field (E_{acc}) at 500 MHz sc cavities first in the world. However, we had no answer that it was surely due to EP. This expectation on EP has been also discussed for a long time among laboratories [1, 2], however, nobody has a clear answer yet. Recently niobium material has been well improved and becomes very reliable. It is the time to come back to this question.

During the last SRF workshop at Saclay and the following TESLA meeting at DESY, "European Headache" was developed: a mystery that KEK's cavities out of the niobium material with $RRR \approx 200$ and treated with EP can achieve high gradients of $E_{acc} > 30$ MV/m [3], but Saclay's ones with $RRR \approx 300$ and treated with CP can not get such a field, and a 1400 °C heat treatment makes possible the high gradients [4]. What is the difference: material, fabrication process or chemistry? For these two years, we had a systematic study intensively using many L-band virgin single cell cavities to get the answer. As a result, we identified "European headache" is owing to CP, and found out the clear superiority of EP over CP for high gradients over 25 MV/m. The results of the experiments are presented in this paper.

2. Experiment

We use many single cell cavities in this experiment. In later sections, we discuss the result with cavity name. At the first we summarize the cavity information in Table 1. Geometrical and RF parameters of cavities are listed in Table 2.

Table 1. Cavity information. Refer to Table 2 on the cavity parameters.

Cavity name	Material	RRR	Fabrication	Cavity parameter	Comments
C-1	Fansteel	350	Jefferson Lab.	A	3.2 mm thick
C-3	Fansteel	350	Jefferson Lab.	A	3.2 mm thick
K-3	Tokyo Denkai	200	KEK	C	2.5 mm thick
K-4	Tokyo Denkai	200	KEK	C	„
K-6	Tokyo Denkai	200	KEK	A	„
K-7	Tokyo Denkai	200	KEK	A	„
K-8	Tokyo Denkai	200	KEK	A	„
K-9	Tokyo Denkai	200	KEK	A	„
K-10	Tokyo Denkai	200	KEK	A	„
K-11	Tokyo Denkai	200	KEK	A	„
K-12	Tokyo Denkai	200	KEK	A	„
K-14	Tokyo Denkai	200-> 400	KEK	B	1400 °C annealed.
K-15	Tokyo Denkai	200	KEK	A	sent Jefferson Lab.
K-16	Tokyo Denkai	200	KEK	A	sent Jefferson Lab.
K-17	Tokyo Denkai	200	Jefferson Lab. (EBW)	A	sent half cells to Jefferson Lab.
N-2	Tokyo Denkai	200	Nomura Plating	A	EBW failure at Iris.
T-1	Tokyo Denkai	200	TOSHIBA	A	EBW failure at Iris.
SHI-1	Tokyo Denkai	200	Sumitomo Heavy Industry	B	
ME-1	Tokyo Denkai	200	Mitsubishi Electric Co. Ltd.	A	ECB

Table 2. Geometrical parameters of cavities used in this experiment.

	A (spherical)	B (spherical)	C (Asymmetric)
R/Q [Ω]	102	110	101
Γ (geometrical factor)	274	266	251.9
Ep/Eacc	1.78	1.89	1.83
Hp/Eacc [Oe/MV/m]	43.8	43.2	45.2
Eacc/SQR(P·Q)	87.35	90.57	86.63
Beam tube diameter [mm]	80	76	80
Inner surface area [cm ²]	1664	1575	1569
A weight per 1 μ m material removal [g]	1.42	1.35	1.34

2-1. KEK in-house fabrication

Here, we present a fabrication procedure of the KEK in-house cavities because most of the cavities used in this experiment were fabricated in KEK [5]. Niobium material is from Tokyo Denkai [6]. Its RRR value is about 200. A half cell is formed by deep drawing with a 80 ton press machine from a sheet of 280 mm in diameter and 2.5 mm thick. After trimming, the whole surface of the half cell is chemically etched slightly for cleaning before electron beam welding (EBW). EBW of equator section of a cell is done from inside with defocus beam. For beam tubes, a sheet is rolled and welded with EBW, then cut into halves, and a mandrel is passed through it to make the real circle. After welding a niobium flange at the end, it is joined with EBW to the iris of the cell.

2-2. Surface treatments

1) Chemical polishing (CP)

Our CP method is a standard buffered chemical polishing. Acid is a mixture of 1:1:1 by HF, HNO₃ and H₃PO₄. We used a fresh acid for every chemistry. Acid temperature is not controlled during CP but usually in between 15 and 45 °C. Just inside of the cavity is removed. The typical material removal speed is 10 μm/min at room temperature. The removed thickness is measured with an ultrasonic thickness meter to check the removal distribution. Its typical material removal distribution is shown in Fig.1 together with the results of EP and BP. These data are for the average material removal of 80 μm calculated from weight difference before and after the treatment. A feature of the removal distribution with CP is a deeper removal at the iris section. It is owing to the upright position of the chemistry. For the other region, it is rather uniform. The material removal rate at the equator section is 90 % of the average removal. Boundary etching for the heavy material removal (≈ 100 μm) might be a problem of CP as discussed later [1]. The typical finished roughness is 5 μm. A rinsing with ultrapure water is carried out immediately after CP, then a high pressure rinsing (HPR) is done for about 1.5 hours with filtered (0.2 μm) demineralized water (10 MΩcm), and finally shower rinsing is done for about 10 min. with filtered (0.1 μm) demineralized water.

2) Barrel polishing (BP)

Barrel polishing (also called tumbling) had been once applied to niobium cavities at DESY [7], but recently KEK also independently developed successfully it [8]. At KEK, plastic chips binding ceramic powder and liquid soap are put in a cavity, then it is rotated horizontally at 100 rpm. The material removal speed is very slow, but this method is very effective to remove irregularities like scratches, or sputter balls by EBW failure [9]. The typical material removal speed is 7 μm/day. It has the largest removal rate at equator section as seen in Fig.1, and it is 250 % of the average removal. This is convenient to grind the EBW seam at equator where the surface should be defect free as the high surface current crosses here. In the TRISTAN-SC project, we polished up the inner surface of every half cell with buffing to initialize the state of surface, then had EBW to complete a cavity. Welding defects were worried about, so every EBW seam was ground mechanically after EBW. BP can finish the both processes at once, so reduces the preparation cost. A problem is embedding abrasives in the polished surface as same as buffing. A light CP (10 μm) or EP (10 μm) is necessary to remove this contamination layer. The finished roughness by BP is 1-2 μm in Rz.

3) Electropolishing (EP)

This is a main interest in this paper. The method is the same as TRISTAN-SCC, namely the horizontal rotating method under a continuous current [10]. Its typical material removal speed is 0.4 μm/min. It has a maximum removal at beam tubes. This is owing to the geometry of electrodes in EP. The equator section is removed by 60 % of the average removal. The finished roughness depends strongly on the initial roughness and the amount of the material removal. Its typical number is 1 μm for the a heavy material removal (≈ 100 μm). EP has less boundary etching as long as EP condition is proper. A problem is a Q-disease owing to hydrogen picked up during EP. We take a medium

temperature (800 - 700 °C) annealing to prevent it. By a recent study, the annealing temperature could be decreased to 600 °C or might be possible even to 500 °C [11]. If taken the annealing and the material of $RRR \leq 200$ is used, the Q-disease does not appear even by an additional heavy EP (130 μm). Rinsing procedure is the same as the case of CP. The hydrogen peroxide (H_2O_2) rinsing used in the TRISTAN SC project was not used in this experiment.

4) High pressure rinsing (HPR)

HPR was done for 1.5 hr in every chemistry as a main rinsing process. The amount of used water is about 800 l. Pressure was initially 85 kgf/cm^2 but decreased to 60 kgf/cm^2 during the frequent uses. This pressure decreasing is mainly due to exhaustion of the pump system. One should notice the chipping of nozzle holes. Actually we found many marks of the chipping on the hole edge by inspection with microscope. This might cause field emission or quench.

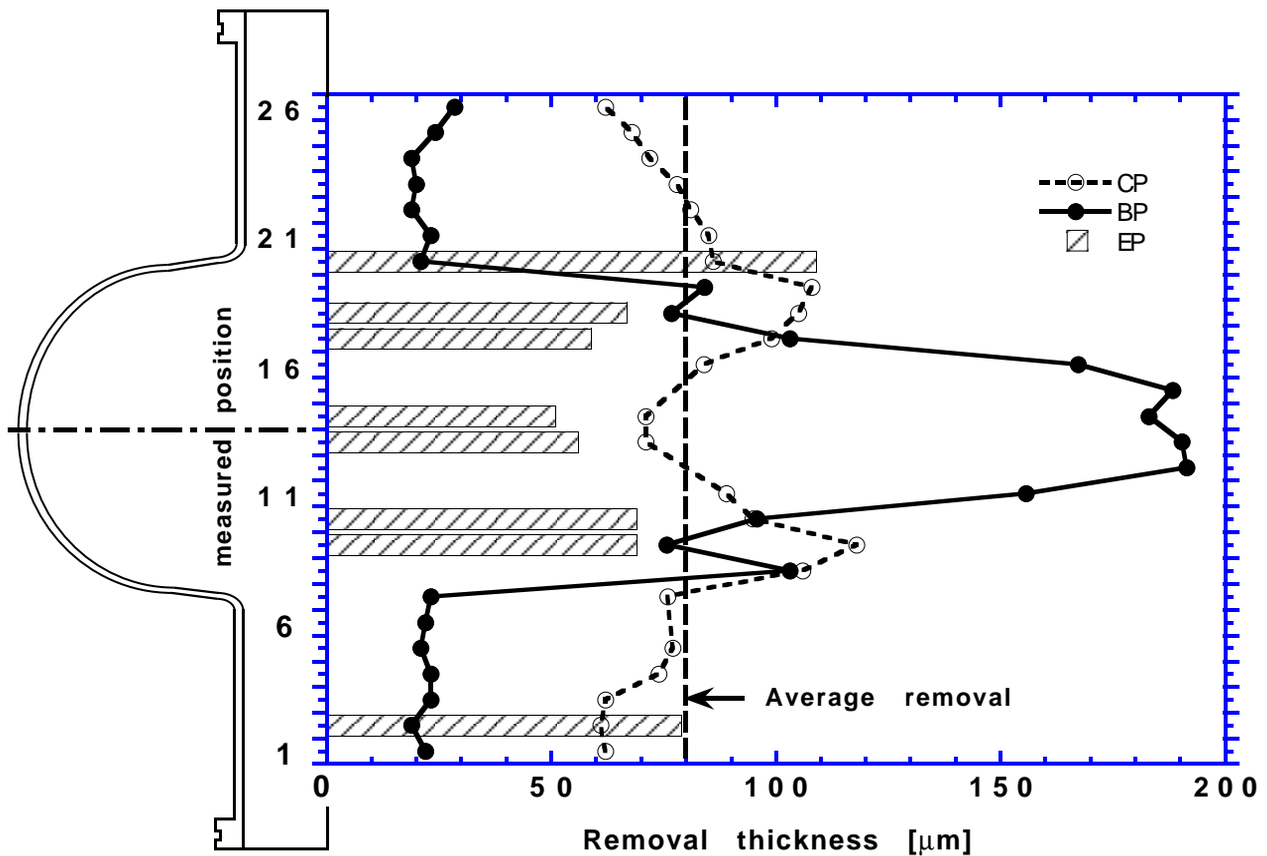


Fig.1 Distribution of material removal with CP, EP and BP. The distribution is for the average material removal of 80 μm calculated from weight difference in each method.

In some cases of field emission, we are suspecting the quality of pure water. In these experiments we used demineralized water (ion-exchanged water) mainly in HPR. At the initial stage in this experiment, there were several cases in which resistivity of water suddenly became worse on the way of HPR. Now the maintenance method for the pure water making system has been well established and such a problem was eliminated. In the following discussion, the experimental results which show heavy field emission at low field less than 10 MV/m are excluded. These are thought due to some accidental

cause such as : nozzle chipping in HPR, water quality, or a cold leak in the cold measurement mentioned later. In such a case, the cavity was just re-rinsed with HPR and re-measured.

2-3. Cold Measurement

After the surface treatment at the company (Nomura Plating), the cavity is immediately brought to KEK and assembled in the class 10 clean room. Indium is used for vacuum sealing of flanges. For vacuum evacuation we have three stands. Each stand consists of a rotary pump (200 l/sec) and a turbo-molecular pump (50 l/sec) as pre-evacuation, and an ion pump (60 l/sec). Cavities are baked around 90 °C for 1-2 days during the pre-evacuation . The typical vacuum pressure prior to the cold test is 1×10^{-9} Torr.

For cold tests, we have two cryostats, one has been used since 1991 [12] and the other was newly built for quick measurement [13]. Vacuum evacuation is continued during cold measurement for the old one, but cavities are sealed with a metal valve for the new one. The pre-cooling has been carried out with liquid nitrogen for one night for the old cryostat, but it is not needed for the new one because the heat capacity is so small. Cooling down speed is about 45 minutes from 300 K to 4.2 K with the new cryostat. The old cryostat can cool down cavities to 1.7 K at best, but the new one is possible to 1.2 K. The old one is a little bit poor with magnetic shielding so that a surface residual resistance of about 3 nΩ adds to the intrinsic one of cavities. The old one needs much time to make it ready for measurement but is reliable. Comparing measurement results with K-14, no difference was seen in both systems. In addition, considering that an $E_{acc} = 39$ MV/m has been achieved with the new system, the blind vacuum sealing during cold measurement is not a problem on cavity performance. In the cases of heavy field emission, vacuum leaking was occasionally found in the new cryostat. Since such a trouble could be a cause of field emission, careful handling is required for the new cryostat. Cavities were cold tested around 1.8 K but some cases were measured at the lower temperature.

3. Experimental Results

Hereafter, we often refer to the amount of the material removal in various chemistries. We define it first. As long as we do not mention especially, an average value for the whole inner surface including beam pipes is used, which is calculated from the surface area and a weight difference before and after the chemistry .

3-1. "European Headache"

So far we have not made systematic investigation on CP with cavity performance. Adding the material removal with CP, we measured the cavity performance to see the "European Headache". We used three virgin cavities (K-7, 10, 11) in order to make sure the reproducibility of the experimental result. BP was not performed before CP for an effect of mechanical grinding not to mix into the result. The chemical material removal was started from 100 μm, then added 30, 50 and 70 μm, up to 250 μm in total where cavity performance is RF considered to be saturated [14]. Cavities were heat treated (750 °C x 5 hr) after the first CP (100 μm) to make it sure that the results are free from the problem of the hydrogen Q-disease. The results are summarized in Fig.2, where the difference of Q_0 value between the measurements is mainly owing to the measuring temperature and the used cryostat. The maximum gradients scatter between 18 to 26.5 MV/m. What is interesting is that not only in the electron loading cases but also in the other cases a steep or exponential Q degradation is observed at high gradients. This new type of Q degradation is also observed in other laboratories with CP cavities [15, 16]. Sometimes we observed this Q degradation with EP cavities but at higher gradients > 25 MV/m. Anyway with CP, the gradient is limited below 30 MV/m either by electron loading or the new Q degradation. " European Headache " has been observed at KEK too.

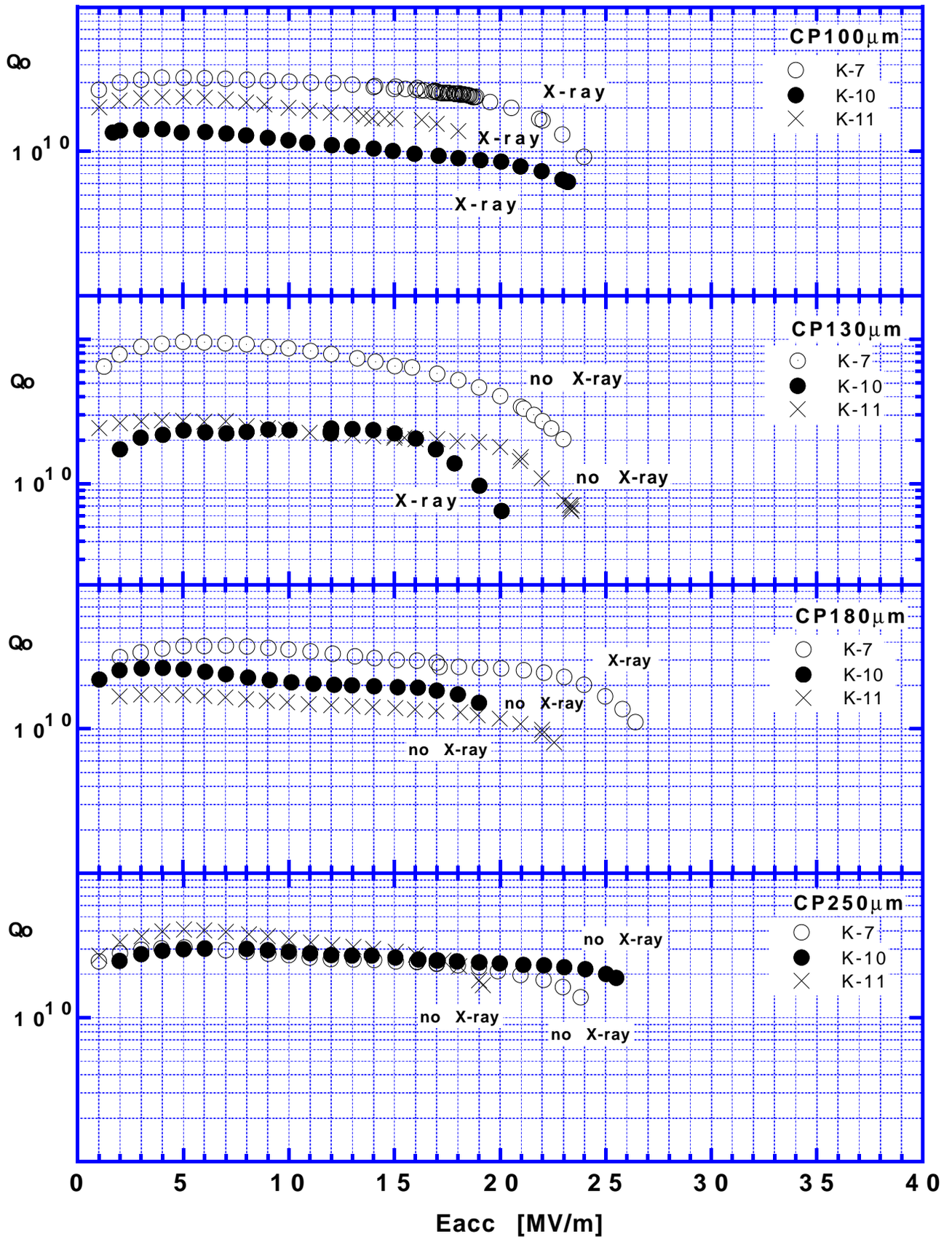


Fig. 2. Cavity performance on chemically polished cavities.

3-2. Effect of light EP on the CP surface

What is the cause of the "European Headache" ? An influence of the hydrogen doped during heavy chemical etching was suspected first. We degassed K-11 at 750 °C again, then chemically polished by 30 μm, however, Eacc,max was not changed: Eacc,max = 22.7 MV/m. That means the contribution of hydrogen is less on the "European Headache".

As the next step, we performed a light EP on these cavities (K-7, 10, 11). The results are shown in Fig.3. K-7 was removed directly 30 μm, then Eacc,max was upgraded to 35 MV/m without any Q-degradation. To see the effect of the adding material removal with EP, K-10 and K-11 were additionally removed every 10 μm, up to 30 μm. At the first 10 μm EP, the gradient was limited at 23.4 (K-10) and 24.4 MV/m (K-11) respectively but the Q-degradation seen in CP was not observed. The additional removals improved Eacc,max, then 30.9 (K-10) and 28.7 MV/m (K-11) were achieved after the 30 μm removal. This excellent effect of EP seems to be owing to not the removal of material defects but the polishing mechanism of EP. This discussion is based on the

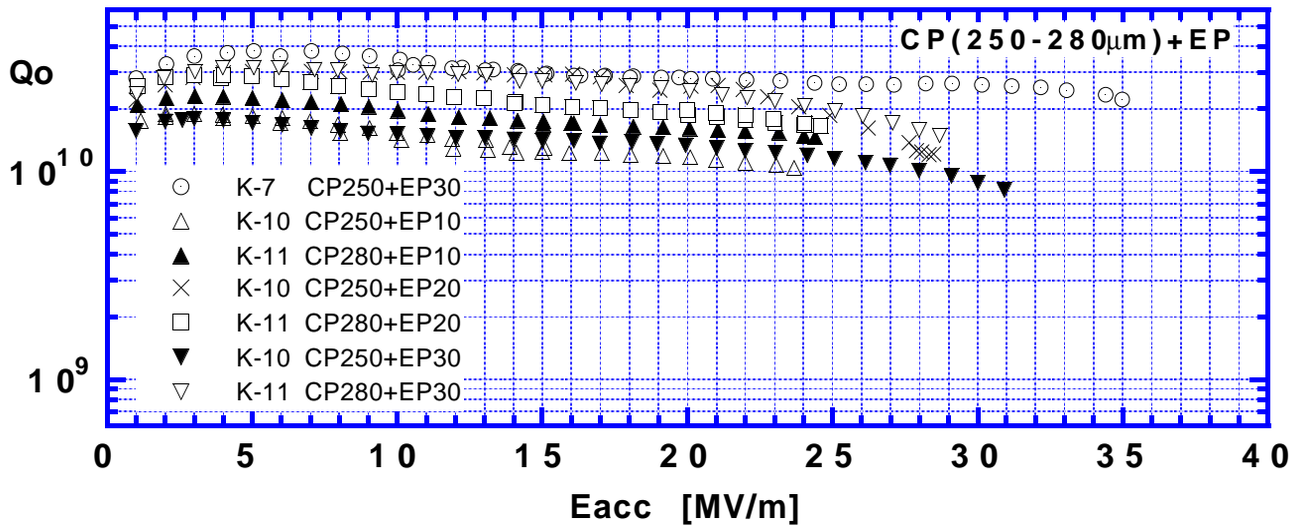


Fig.3 Effect of the light final EP on chemically polished cavities.

hypothesis of defects free material [17] but if not so, such a consistent result could not be obtained in many cavities. From these experimental results, it is pointed out that the "European Headache" has the cause in chemical polishing itself. This will be discussed more in section 4. Here, we can see a superiority of EP over CP.

3-3. Chemistries and cavity performance

As mentioned initially, now we have three kinds of polishing techniques: CP, EP and BP. CP has a very fast material removal speed but seems to be unacceptable for the high gradients Eacc > 30 MV/m as seen in the "European Headache". If taking a high temperature annealing, it is provably possible but other problems will occur: softening in material, cost increasing so on. EP has a rather slow removal speed but offers the excellent cavity performance. BP is a very reliable grinding method for irregularities but has a contamination problem of abrasives. To understand the mechanism of the field limitation, let's see the correlation between the chemistry and Eacc,max for various chemistries. And such an investigation will show more clearly the superiority of EP.

(1) EP + HPR

A question with the material removal has been made clear with CP by Dr. Kneisel [14, 16]. Comparing EP and CP on this subject is very interesting to study the characteristics of EP. At first, we made the experiment to see the improvement in $E_{acc,max}$ by adding material removal with EP. BP was not taken before EP for its effect not to mix into the result. The SHI-1 cavity was chemically polished at first by 20 μm to remove the surface contamination. Then annealing (800 - 700 x 5 hr) was performed to avoid the hydrogen Q-disease problem. Finally it was electropolished every 30 μm , up to 120 μm in total. The results are presented in Fig.4. In the first measurement, field emission was

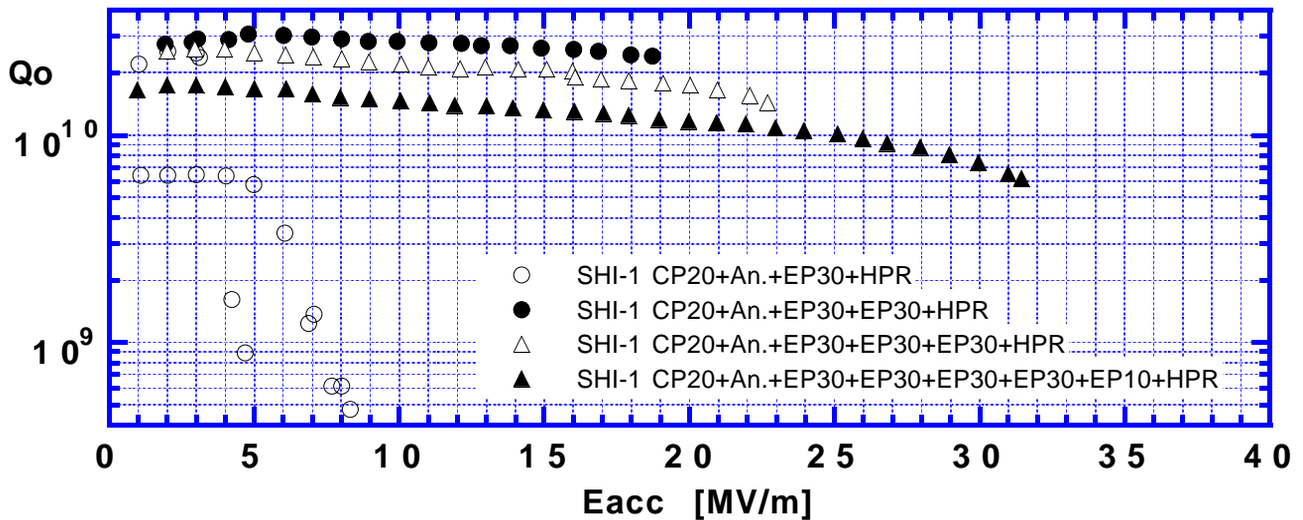


Fig.4 Improvement of cavity performance by the additional EP material removals.

observed. The cause was not searched enough and not clear. On the 4th trial (120 μm EP), we met a vacuum leak accident, so the measurement was not performed. The final measurement was done after an additional 10 μm EP to remove the contamination. The $E_{acc,max}$ improved gradually by the additional removals and finally achieved 31.5 MV/m after 130 μm EP. The detail information with this experiment is reported in the reference [18].

(2) BP + EP(light) + Annealing + HPR

BP is very reliable to remove surface irregularities [9]. That could offer a good quality control on the cavity performance. It is easy and cheap. To search a surface treatment method very reliable in the cavity performance and low cost in the preparation, it is worth to investigate the procedure which consists of mainly barrel polishing, and light EP to remove the contaminated layer by BP. We performed the experiment using five virgin cavities (K- 4, 8, 9, 12, N-2). Even lightly, the EP after BP makes the hydrogen Q-disease. In this experiment, a medium temperature annealing is needed to prevent this problem. After the annealing, cavities were just rinsed by HPR with demineralized water (10 M Ωcm). The results are shown in Fig.5. We used a light EP (10 μm) or a combination of CP (5 - 7 μm) and EP (10 - 30 μm) to remove the contaminated layer by BP, which were considered to be enough to eliminate the abrasive contamination. K-4 achieved 33.1 MV/m in $E_{acc,max}$ but the others were limited around 25 MV/m.

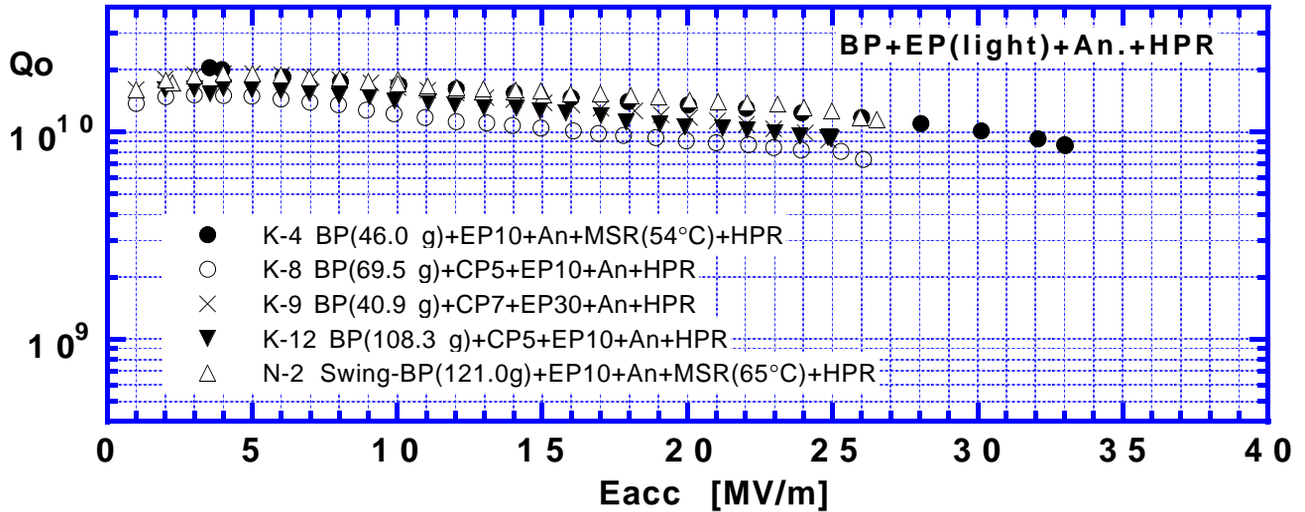


Fig.5 Cavity performance with the procedure: BP+EP(light) + Annealing (750 °C) + HPR.

We tried megasonic rinsing (MSR) for K-4 and N-2 prior to HPR in order to strengthen rinsing process, however, the excellent result in K-4 is still not sure due to this effect or not. The barrier at 25 MV/m might have a mechanism behind. It will be discussed later. K-12 had many sputter balls around EBW seam on the equator due to the failure of EBW, and N-2 also had an EBW failure at the iris section, but the same results were obtained by this procedure with more removal by BP.

(3) Effect of the final light EP after annealing

To try to improve the maximum gradients and to see the effect of additional EP, three cavities (K-8, 9, 12) were electropolished with a 10 μm step up to 30 μm. The results were presented in Fig.6.

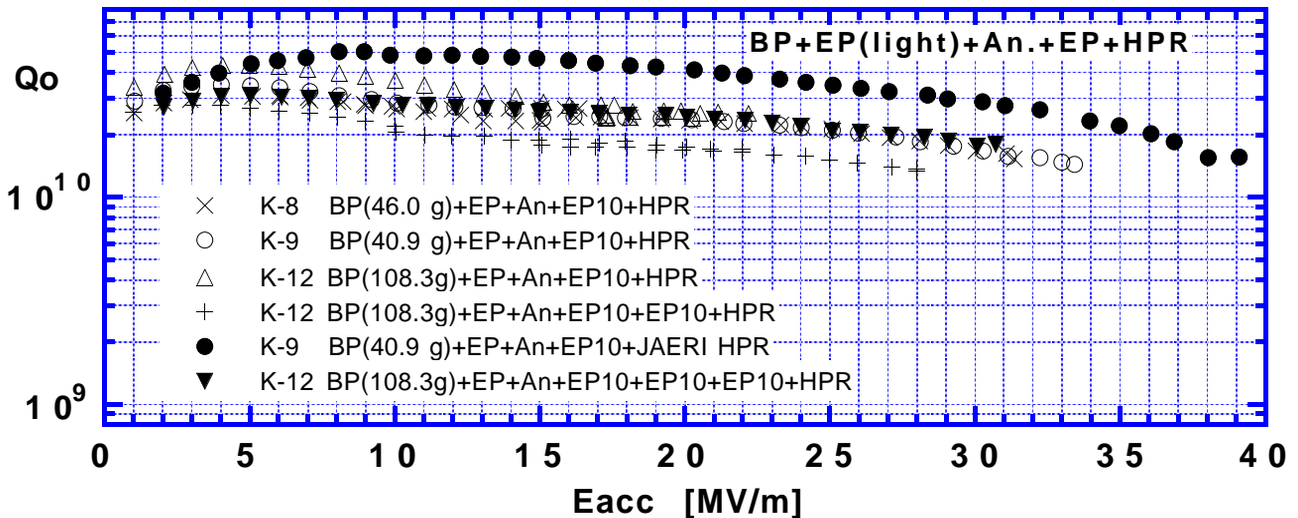


Fig.6 Cavity performance with the procedure: BP+EP(light) + Annealing (750 °C)+ EP (10-30 μm)+ HPR.

For K-8 and K-9, the gradient upgraded without any Q-degradation to 31.4 (K-8) and 33.4 MV/m (K-9) by the first 10 μm finishing. With K-12, Eacc,max improved to 29.3 MV/m by the first 10 μm EP and up to 30.7 MV/m finally by 30 μm EP. The final light EP is very effective to eliminate the barrier at 25 MV/m in the procedure (2).

After this experiment, K-9 was just rinsed again with HPR at JAERI which was newly built. Eacc,max reached to 39 MV/m at 1.5 K (●) but 31.5 MV/m at 1.8 K. Eacc,max strongly depended on the liquid helium temperature. This cavity was left in the class 100 clean room for several months just covered with aluminum foils since the latest cold test. We are not sure whether this phenomenon depends on the rinsing or not, but this is a very interesting phenomenon as a future investigation.

(4) Heavy EP after barrel polishing

We confirmed the final light EP can eliminate the barrier at 25 MV/m in the procedure (2), however, a question still remains for the cause: an influence of BP or the contamination during annealing. To get the answer, we took a heavy EP instead of a light EP in the procedure (2). Two virgin cavities (K-3, T-1) were used in this experiment. The result is shown in Fig.7. A Q-degradation with x-ray (field emission) was observed clearly at high gradient with K-3, but the Eacc,max was achieved over than 30 MV/m in both cases. Combining this result and one of (3), it is suggested that the barrier at 25 MV/m is not owing to the contamination by annealing but the influence of BP, and it might be still remained in the final light EP of about 30 μm . The abrasive contamination by BP is removed by 10 μm material removal with EP or CP. Therefore, a following picture is imaged for the barrier that BP makes damages on the surface, its influence is extended underneath about 30 μm , and it is not recovered enough by the medium temperature annealing (700-800°C x 5 hr). The Q-degradation, which is due to field emission mainly or partly, might be owing to the contamination by annealing or the limitation in our rinsing technique.

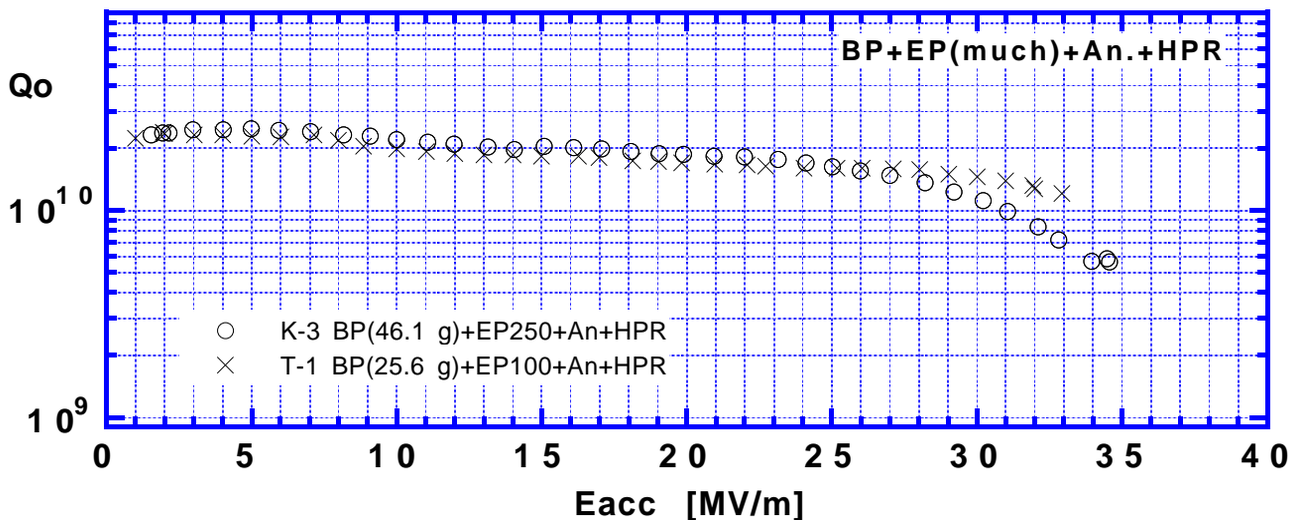


Fig.7 Cavity performance with the procedure: BP+EP(heavy)+ Annealing (750 °C) + HPR.

(5) Superiority of EP over CP

Here, we change EP with CP in the treatments (2)-(3) to see the polishing effect of CP. Results are shown in Fig.8 and Fig.9. Fig.8 is corresponding to the case that EP was changed with CP in the

procedure (2). This experiment was done with the N-2 cavity and repeated twice to confirm the reproducibility. As shown in Fig.5, this cavity had achieved 26 MV/m by the treatment (2) before these tests. In this tests, the amount of BP is not much because its purpose is to just reset the surface to the mechanically ground state. Here, we tried megasonic rinsing (MSR) prior to HPR in the rinsing procedure but it would not bring any confusion to the explanation. The result on the N-2 cavity in the treatment (2) was obtained by the material removal of 10 μm by EP after BP. In this experiment, 10 or 15 μm was removed by CP. It should be enough for the high gradient of 25 MV/m as the material removal. However, as seen in Fig.8, Q-degradation without x-ray appeared and limited the gradient to 22 and 24 MV/m. This Q-degradation is common to CP, remembering the result of the "European Headache".

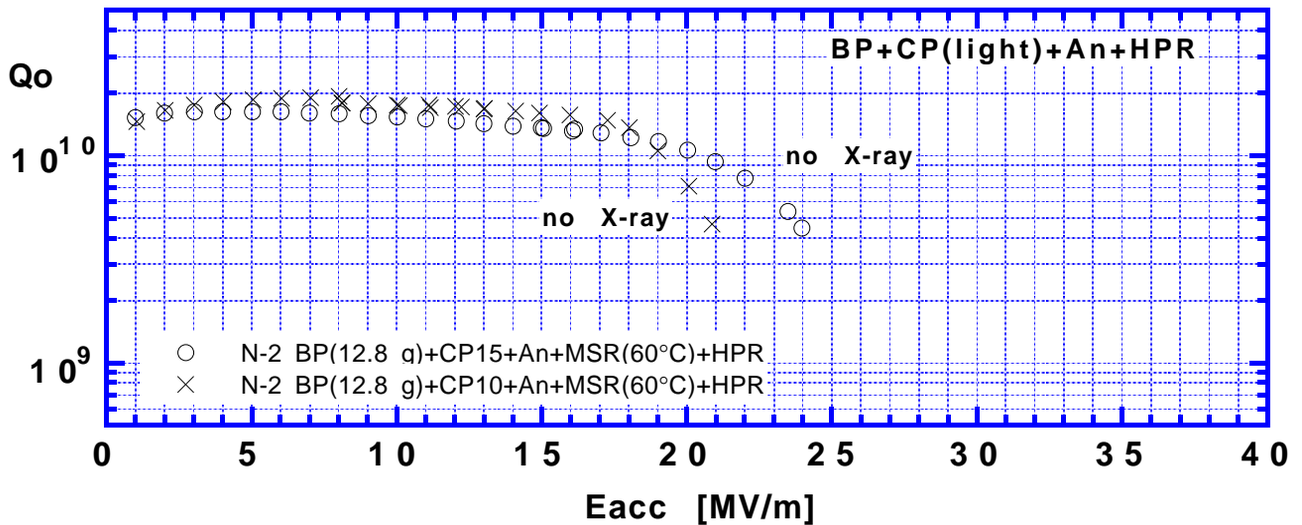


Fig. 8 Effect of a light final CP on cavity performance.

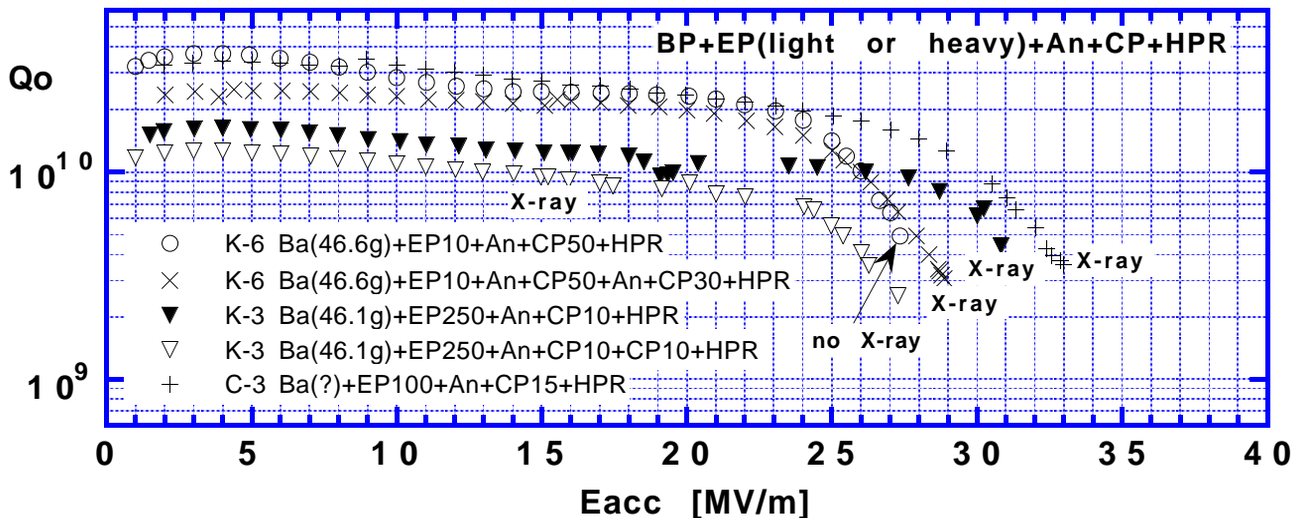


Fig. 9 Deterioration of cavity performance by CP.

The result of K-6 (○, X) in Fig.9 is for the case that the final EP was changed with the final CP in the procedure (3). With EP, the final removal of 30 μm is almost enough for high gradients : $E_{acc,max} \geq 30$ MV/m (see Fig.6). In this experiment, the Q-degradation without x-ray appeared again, and $E_{acc,max}$ was limited to 27.3 MV/m in spite of 50 μm removal.

The results of K-3 (▼, ▽) and C-3 (+) in Fig.9 are for the cases added a light CP after a heavy EP. If an additional CP is really light ($\approx 10\mu\text{m}$), the results are better than those obtained with CP only (compare to Fig.2). But a tendency of performance degradation with an additional CP is seen in the results of the K-3 cavity. Probably CP makes the electropolished good surface worse.

As is shown with many experimental data, the superiority of EP is clear. High gradients (> 30 MV/m) can be achieved by the final material removal of > 30 μm by EP with any heavy pre-polishing. On the other hand, if CP is used as a final finishing, the Q-degradation appears and limits the high gradient. Now it can be said that the "European Headache" is a Q-degradation starting around 15 - 20 MV/m with or without x-ray detection. High temperature heat treatment : HTT(> 1300 °C) or a new rinsing technique is necessary to achieve 30 MV/m with CP.

4. Discussion

A. The dependence of the $E_{acc,max}$ on the material removal thickness

The dependence of the $E_{acc,max}$ on the material removal thickness is very interesting itself and is necessary to be made clear for further discussion or understanding. Fig. 10 shows this plot for our present data and the data from Dr. Kneisel (K-17) [16], which were all obtained with a 1.3 GHz single cell cavity made from Tokyo Denkai material of $RRR \approx 200$. Since we used three different polishing methods and they have different removal distribution, the amount of material removal is evaluated at an equator region where the surface current has the maximum. At a glance, we can point out three characteristics. Firstly, below ≈ 100 μm the maximum gradient increases with a removal thickness. Secondly, above ≈ 100 μm, the maximum gradient scattered in wide range and looks to be saturated. But thirdly, the averaged maximum gradient for cavities finished by EP is higher than that for cavities finished by CP. Necessity of the heavy material removal has been already shown by P. Kneisel, and it suggests the existence of the damage layer of ≈ 100 μm thick, which is probably introduced by forming or EBW. In these cases where cavities are polished heavily (≥ 100 μm) by CP only, the maximum gradient seems to be limited by the surface condition peculiar to CP. On the other hand, in the EP finished case, it is not clear whether the finishing by EP of about 30 μm is enough or the surface condition made by CP or BP still remains, or our rinsing and assembling technique limits the maximum gradient.

The dependence of surface roughness on removed thickness for CP and EP in Fig.11, 12 respectively might be helpful for further understanding. The data were taken using BP samples. In the case of CP, saturation is seen above 100 - 150 μm, but the saturated state seems to depend on the initial state. On the other hand, EP gives almost the same saturated state but the necessary material removal depends on the initial roughness. So, in the case of CP + EP in Fig.10, the EP of ≈ 30 μm might be not enough, because the roughness after heavy CP is 2 - 5 μm. However, the maximum gradient of the cavities, which are finished by EP with a smooth surface by BP (≈ 1 μm), or ECB : Electro-chemical buffing (Fig.13), which is reported in reference [19], is also limited to the same level. Above discussion might indicate that the surface finished by EP is already smooth enough but the rinsing or assembling technique limits the maximum gradients. Further experiments are necessary for the full understanding of the effect of polishing mechanism.

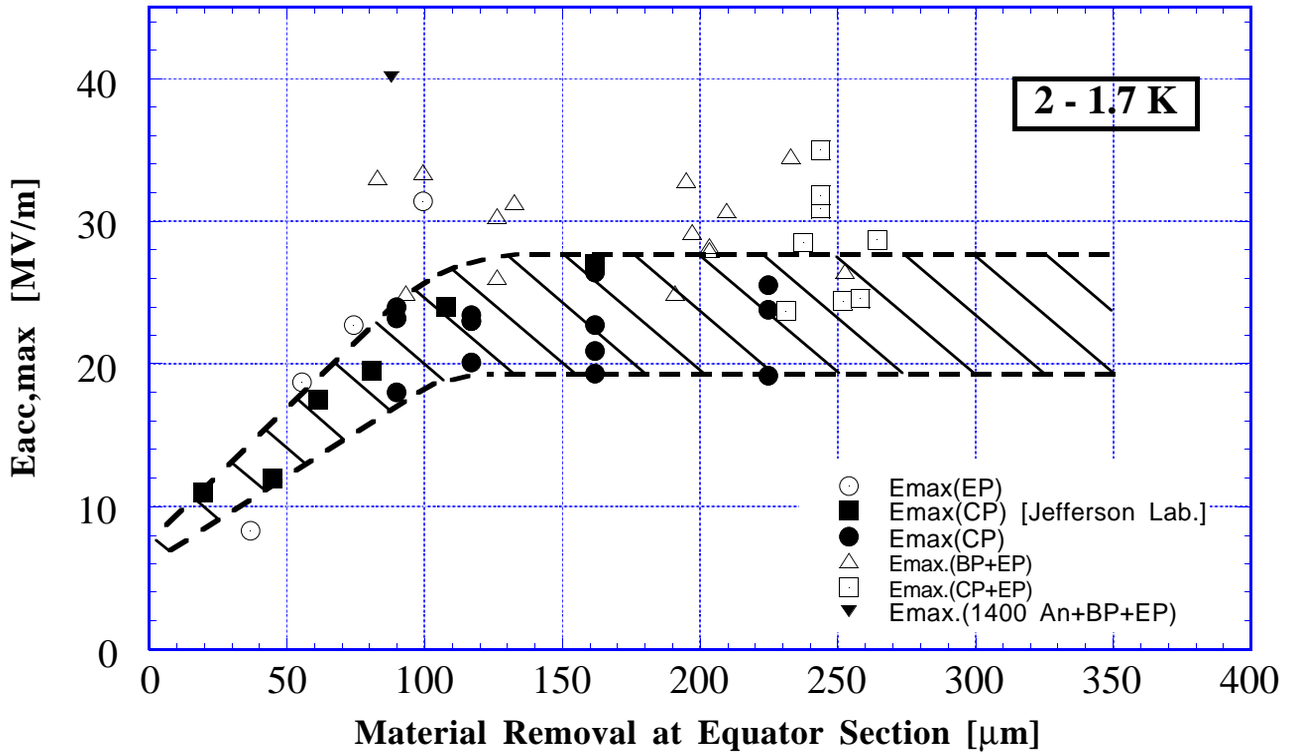


Fig.10 Relationship between a material removal at equator section and the maximum field gradient.

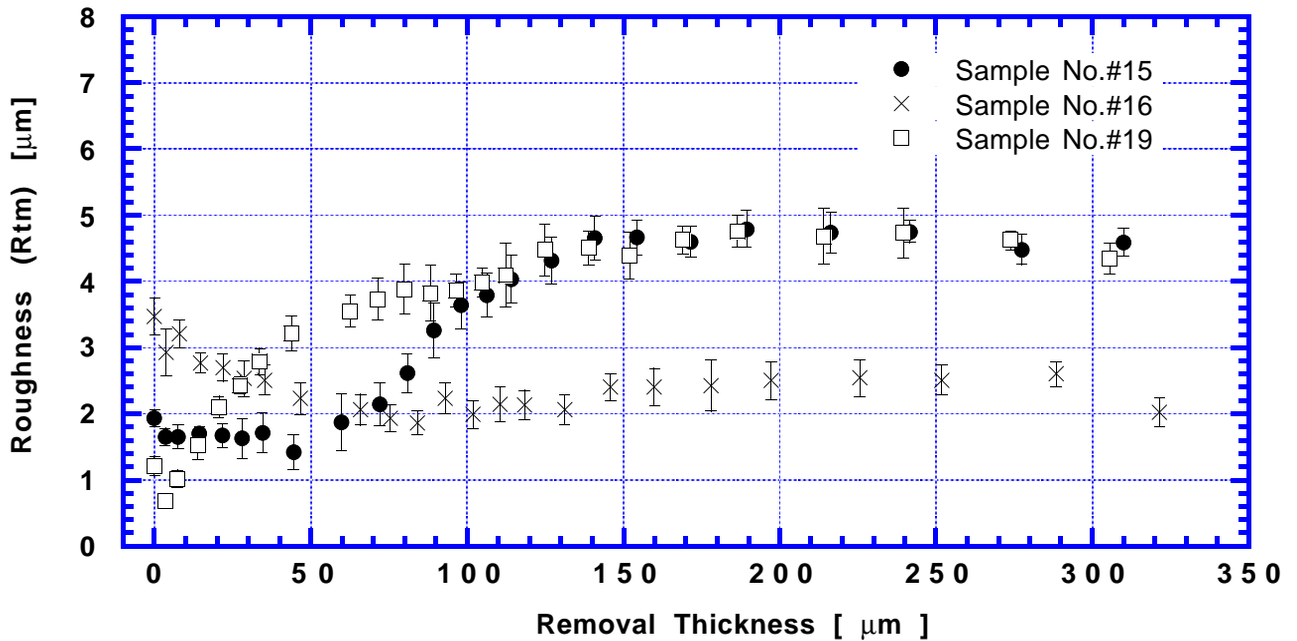


Fig.11 Relationship between material removal and surface roughness with CP. *The error bar means the variation in three points measurements.*

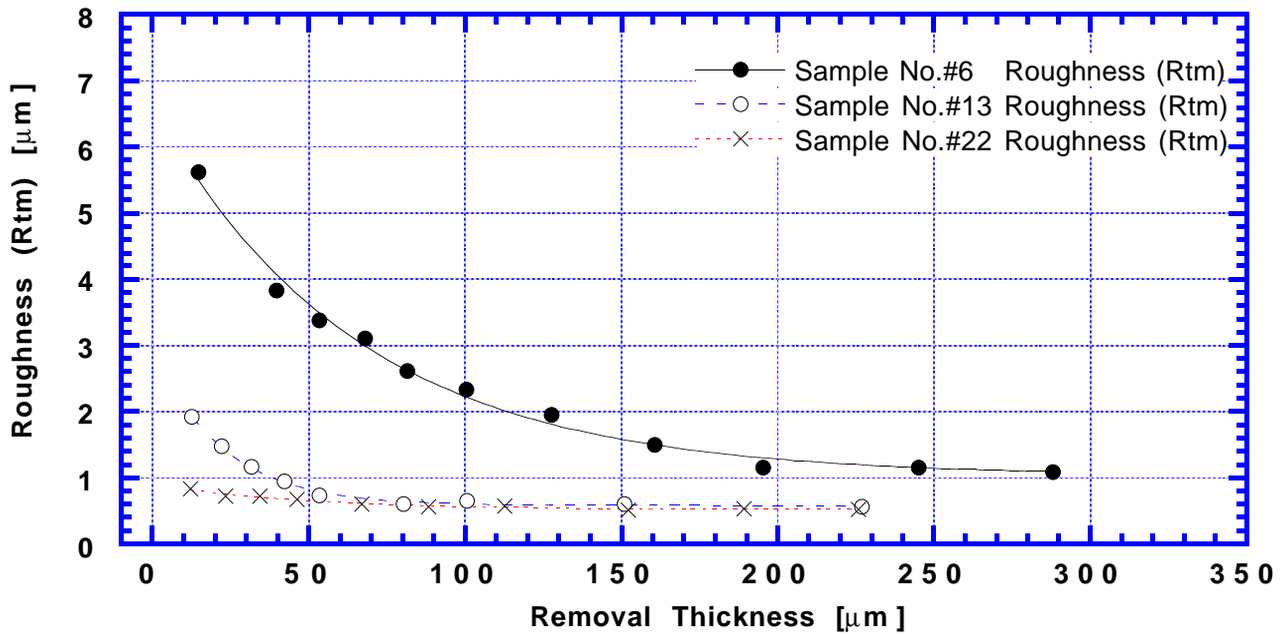


Fig. 12 Relationship between material removal and surface roughness with EP.

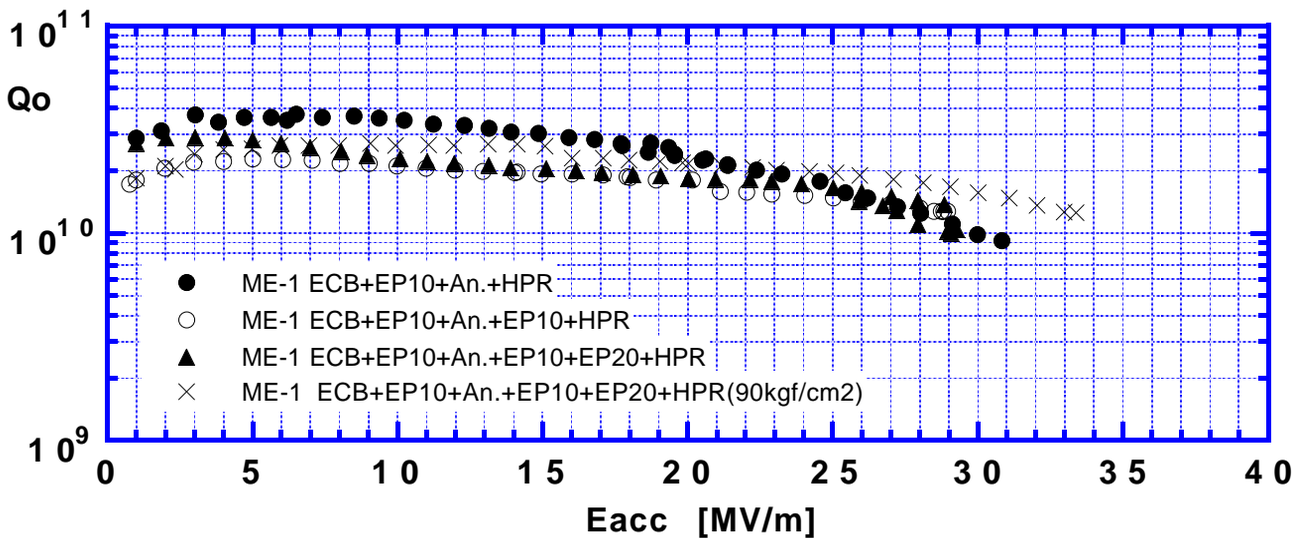


Fig.13 Cavity performance with a smoother surface by ECB.

B. Cause of the "European Headache"

It is well known that CP etches crystal boundary much more than EP. By comparing grain size (typically 50 - 100 μm) of the samples and the structure of output signal of the surface roughness meter, we can consider that the saturated surface roughness after heavy CP ($\geq 100 \mu\text{m}$) mainly comes from etched crystal boundaries. So, it is very natural to search a cause of the "European Headache" in the etched boundary. Surface damage like as cleavage, which can be made by stress concentration by forming, concentrates at grain boundaries. Dislocation generated during the fabrication process (forming or EBW), or segregation of impurities also is concentrated there. Grain

boundary is very unstable in energy and reacts actively with acid, then results in boundary etching in CP. As boundary etching going ahead, grain boundary edge is grown, then chemical residue or particle contamination become easier to remain there. It would be harder for grain boundaries to be cleaned up even using HPR. Furthermore, hydrogen cracking or formation of niobium hydride, which is a weak superconductor, might happen at the boundary by such an active chemical reaction. These could be a cause of heating spot. Namely, two characteristics of the boundary etching, the geometry and the chemical products, could be pointed out as the cause of the Q-degradation in CP cavities. HTT ($\geq 1300\text{ }^{\circ}\text{C}$) makes the crystal size larger, also recovering of damaged surface and the evaporation of impurities at the boundary are expected. As a result, the density of boundary becomes lower and the boundary etching might become lighter in CP. Thus, HTT can achieve the higher maximum gradient without the Q-degradation (see Fig.14), though the improvement of thermal conductivity of niobium material by HTT also would play an important role. Thus, the boundary etching hypothesis could explain the Q-degradation seen in the "European Headache" and the upgrade of the gradient in the CP cavities after HTT.

Boundary etching would strongly depends on the state of surface damage. Like BP, where many damages are introduced, it would become heavier. This might be a reason why BP+ light CP did not work well in Fig.8. The boundary etching hypothesis indicates difference in the Q-degradation in the "European Headache" with forming method. As a future study, comparison the Q-degradation in CP among deep drawing, spinning or hydro-baldge forming, and annealing prior to CP is very interesting. One reason why KEK also observed the " European Headache " might be that Saclay and KEK used the same forming method : deep drawing.

The boundary etching hypothesis also calls attention to the HTT with double sided titanium. In such a case, titanium might precipitate much or might be concentrated around crystal boundaries [20]. This titanium concentration would be difficult to be eliminated, or would be promote the boundary etching, resulting in the Q-degradation at the high gradient. At Saclay after HTT with double sided

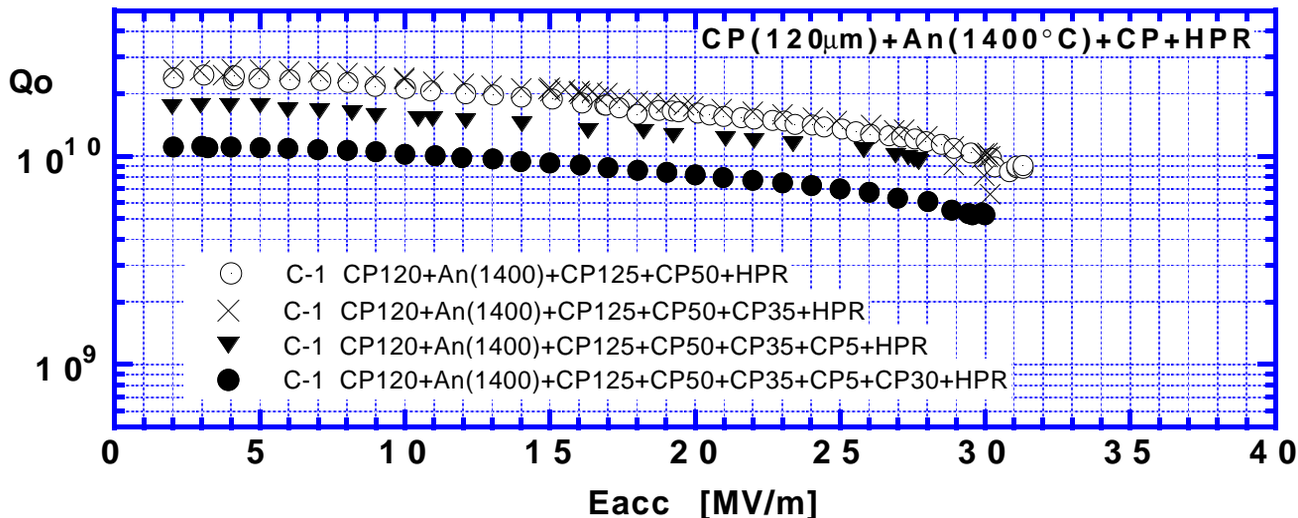


Fig.14 Cavity performance by HTT with single sided titanium.

titanium, still Q-degradation is observed in CP cavities. KEK had also observed the Q-degradation (heavy field emission) on a cavity (M-2: not used in this experiment) with 60 μm CP after HTT with double sided titanium [12]. On the other hand, a single sided titanium HTT is almost free from these problem. This might be a reason why the C-1 cavity treated with this method achieved the high gradient $\approx 30\text{ MV/m}$ without the Q-degradation (Fig.14).

"European Headache" might be solved by a new rinsing method. Really the world record is obtained with a CP cavity (JL-1) made from Tokyo Denkai RRR ≈ 200 at Jefferson Lab. [17], which is our mystery : "Japanese Headache". However, this does not deny the superiority of EP, since the improvement of the maximum gradient can be expected also for EP cavities with the new rinsing method. The JL-1 cavity is now at KEK in order to confirm the superiority of EP. In the first measurement at KEK after just rinsed with HPR, $E_{acc,max}$ was limited at 13 MV/m. This cavity is waiting a cold test after EP. The further experiment is under going.

C. Processing levels at KEK

We have a problem to overcome with our cavities. We observe often processing levels accompanied x-ray at $E_{acc} = 15 - 20$ MV/m and around 25 MV/m. In many cases these levels are easily processed out, but sometimes partially processed out or can not be overcome. And in some cases there is nothing at all. It has no correlation with chemistry, EP or CP. Changing the demineralized water in the rinsing to ultrapure water, the situation was not changed. Oil free evacuation was tried with a scroll pump for rough pumping instead of the rotary pump, but there was no change. Alcohol rinsing was also tried but the result was the same. The similar but much light processing level is observed at Saclay [21]. It is rare at Jefferson Lab [16] and there is no report from DESY.

We were suspecting multipacting for this phenomenon [22, 23]. Multipacting was reported for the cavities with spherical shape [24, 25]. To make it clear, recently we had an experiment in a collaboration with Jefferson Lab, where the secondary electron yield of the niobium surface was intentionally increased by oil contamination. Processing levels were observed at 14, 17 and 25 MV/m in this experiment. Recently computer calculations have been carried out by Dr. R. Parodi at INFN Genoa for this result and has predicted one point MP at 14 MV/m and two point MP at 25 MV/m [16]. For the level of $E_{acc} = 17$ MV/m, the trajectory simulation has not yet succeeded and further calculations are needed. The barrier at 25 MV/m seen in Fig.5 might be related to the multipacting, however, a further investigation with temperature mapping and x-ray mapping is needed.

5. Possible chemistries for $E_{acc} = 40$ MV/m

For the future superconducting liner colliders like TESLA-1000, upgrading the gradient is very important. From our experiments, we can propose several procedures based on the finishing by EP, as candidates to achieve 40 MV/m. In any case, more powerful rinsing method is desirable or might be indispensable to improve the reliability. If the annealing furnace is clean enough, the final EP can be done before 750 °C annealing.

(1) Heavy Removal ($> 100 \mu\text{m}$) by BP or CP or EP + Annealing (750 °C) + EP($> 30 \mu\text{m}$) + HPR

With these procedures, we have already achieved 35 MV/m without a Q-degradation. The necessary removal thickness is not yet known, and an improvement might be expected by the optimization of the amount of the final EP.

(2) HTT($> 1300 \text{ }^\circ\text{C}$ with Ti) + EP($> 100 \mu\text{m}$) + Annealing(750 °C) + HPR

If HTT ($> 1300 \text{ }^\circ\text{C}$) is available, it will improve the reliability for 40 MV/m. This procedure is the best one we can propose at present. HTT is not necessary to be performed for full cavities. Half cell HTT is preferable, because it does not need a large furnace, very good vacuum and prudent cares for handling, deformation or contamination. The chemistry must be an EP because of the problem of the boundary etching discussed before. As is shown in Fig.15, we have achieved 40 MV/m without a heavy Q-degradation by this procedure essentially. The effect of BP is not clear but the reason of a light BP added is that we had not enough reliability on EBW at that time. The detail of this experiment is reported in the reference [26].

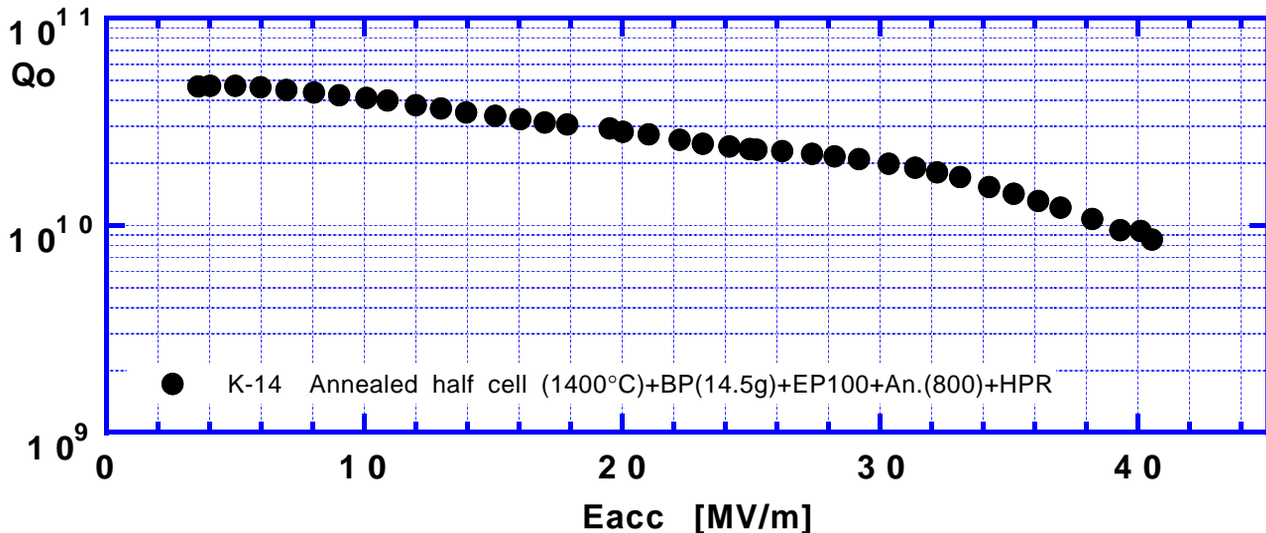


Fig. 15 Cavity performance by the procedure: Half cell annealing (1400 °C with Ti) + BP (14.5 g) + EP (100 μm) + Annealing (750 °C) + HPR.

6. Summaries

By a series of our experiments in the last two years on the 1.3 GHz single cell niobium cavities, the followings are made clear:

- 1) The "European Headache" seen at Saclay is observed also at KEK. This is characterized by a steep Q-degradation with or without x-ray detection, starting at the gradient below 25 MV/m. This phenomenon is common to the chemically polished cavities, so it should be called as " CP Headache ".
- 2) It is very hard to achieve higher gradient > 25 MV/m by the conventional CP. On the other hand, the high gradient over than 30 MV/m can be achieved by the finishing with EP irrespectively of the method of pre-polishing. The superiority of EP over CP is clear.
- 3) The reason why KEK have attained the gradient > 30 MV/m so often is owing to the EP, and not to the material.

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