Aging in the Large CDF Axial Drift Chamber


Abstract—The Central Outer Tracker (COT) is a large axial drift chamber in the Collider Detector at Fermilab operating with a gas mixture that is 50/50 argon/ethane with an admixture of 1.7% isopropanol. In its first two years of operation the COT showed unexpected aging with the worst parts of the chamber experiencing a gain loss of ~50% for an accumulated charge of ~35 mC/cm. By monitoring the pulse height of hits on good tracks, it was possible to determine the gain as a function of time and location in the chamber. In addition, the currents of the high voltage supplies gave another monitor of chamber gain and its dependence on the charge deposition rate. The aging was worse on the exhaust end of the chamber consistent with polymer buildup as the gas flows through the chamber. The distribution in azimuth suggests that aging is enhanced at lower temperatures, but other factors such as gas flow patterns may be involved. Elemental and molecular analysis of the sense wires found a coating that is mostly carbon and hydrogen with a small amount of oxygen; no silicon or other contaminants were identified. High resolution electron microscope pictures of the wire surface show that the coating is smooth with small sub-micron nodules. In the course of working with the chamber gas system, we discovered a small amount of O$_2$ is enough to reverse the aging. Operating the chamber with ~100 ppm of O$_2$ reversed almost two years of gain loss in less than 10 days with an accumulated charge of ~2 mC/cm.

I. INTRODUCTION

The Collider Detector at Fermilab (CDF)[1] is a large magnetic detector built to study 2 TeV PbarP collisions at the Fermilab Tevatron Collider. At its center is the Central Outer Tracker (COT)[2], a large cylindrical tracking chamber coaxial with the beam that measures the 3-momentum of charged particles with excellent resolution. This paper discusses the aging observed in the COT and our efforts to remedy it. Section II reviews the technical details of the chamber. Section III describes the definitive evidence for aging in the chamber. Section IV discusses the nature of the aging seen in the COT and our initial ideas for restoring chamber gain. Section V describes the steps we took and how they led to adding oxygen to reverse the aging in the COT.

Section VI reviews some of the tests and studies we made to get a better understanding of the aging. Section VII presents the conclusions.

II. DETAILS OF THE COT

The COT resides inside a 1.4 T solenoid magnet in the middle of the CDF detector. It is a cylinder that measures approximately 3 m long by 3 m in diameter. There are more than 30,000 sense wires and 40,000 potential wires strung between two thick aluminum endplates. The wires are arranged radially in 8 annular regions called super-layers (4 axial and 4 small-angle stereo). Inside each super-layer the wires are organized in a jet cell configuration with each cell tilted 35º with respect to the radial direction. The variable in the axial direction is $z$ with the chamber center at $z = 0$. The azimuthal variable is $\phi$ which has a correspondence with the super-layer cell number.

There are 12 sense wire layers per cell giving a total of 96 sense wire layers in the 8 super-layers. Between each pair of sense wires and at the end of the cell are “potential” wires that improve the shape of the drift field. The potential wires operate at lower voltage than the sense wires and the electric field at their surface does not generate gas gain. Both sense and potential wires are 40 µm gold plated tungsten. The cathodes are vapor-deposited gold on 6.35 µm mylar foils. The maximum drift distance is 0.9 cm.

The chamber gas is argon/ethane (50/50) with an admixture of 1.7% isopropanol. Initially the gas flow was 20 SCFH which corresponds to one volume exchange every 30 hours. Test monitor chambers saw no signs of aging.

There were several features built into the system to help monitor the gain of the COT. In order to facilitate dE/dx measurements in the data, the amplifier-shaper-discriminator sets its output pulse width proportional to the integrated input charge. The width of all hits, and thereby the pulse-height, are stored in the CDF physics data sets allowing a determination of the gain as a function of position in the chamber. In addition all the high voltages and currents are archived every ~10 minutes. Archives are also kept of the Tevatron luminosity and some of the loss monitor rates.
III. EVIDENCE FOR AGING

Despite the mechanisms in place to monitor chamber gain, detecting aging was not easy. There were hints early in 2003 and through the summer. The online track trigger which has limited redundancy exhibited phi dependent inefficiencies. Definitive evidence of aging came from careful studies of the offline tracking in November-December 2003. They showed that the gain was reduced in the inner super-layers relative to the outer super-layers and the gain was reduced at the exhaust end of the chamber relative to the input end as shown in Fig. 1. The two points in the middle have lower gain because of a wire center support. Also the gain was reduced in some phi regions, especially below the beam as shown in Fig. 2.

In March 2004 an aged wire plane was removed from an inner super-layer of the COT. Scanning Electron Microscope (SEM) measurements were made of a piece of aged wire and compared to a piece of new wire. The surface of the new wire looks brighter and striations are visible (see Fig. 3). The aged wire is dark and smooth except for the sub-micron nodules dotting its surface (see Fig 4).

Electron Dispersive Spectroscopy (EDS) of a new wire showed mostly gold with a little background carbon while the aged wire had a much larger carbon peak and a small oxygen peak (see Fig. 5). EDS uses an electron beam to eject electrons from the inner shells of atoms. The x-ray spectrum generated...
when these states are refilled gives a measure of the elemental
cleanup of the sample. It is not sensitive to hydrogen.

Some additional measurements were made to study
molecular bonds. An XPS scan indicated primarily CC and CH
with a few CO bonds. A Fourier Transform Infrared (FTIR)
analysis of the absorption of reflected broadband infrared
photons also gives a measure of molecular bonds. It showed a
broad absorption band associated with OH bonds and a cluster
of narrow absorption bands associated with CH$_2$ and CH$_3$
bonds.

We found no evidence for aging elsewhere in the COT
super-cell. The EDS spectrum of a potential wire in the
extracted plane was consistent with that of a new wire. The
resistance of the surface of a cathode sheet in that cell was
consistent with a new sheet and there was no visible coating.
We did not extract a cathode sheet, so it was not possible to
perform further microanalysis.

IV. AGING HYPOTHESIS

The aging appears to be due to a coating on the sense wires.
Presumably polymers, strings made up of groups of CH$_2$, grow
in the avalanche environment as the gas moves down the wires.
When the strings become long enough, they plate out on the
wire. Presumably this “condensation” is temperature
dependent. Probably there is also ablation in the avalanches
and aging depends on the balance between ablation and
deposition. No evidence was found for silicon or other
contaminants in the gas that might be causing the aging,
although we can’t rule out contamination from some aliphatic
oil. However, it appears likely that the aging seen in the COT
would be present in most hydrocarbon gases. It is a type of
aging that is not easy to reproduce in a small chamber with a
localized source. The polymers probably have to stay in a
radiation environment a long time in order to become copious
enough and to grow long enough to make significant deposits
on the wires. Also, since it is a multi-step process, the gain
falls faster than linearly with the collected charge on a wire.

In order to combat this sort of aging there are several
straightforward steps that can be taken. Several groups have
found that increasing the gas flow helps. The purpose is to
vent the polymers before they grow long enough to deposit.
Increasing the temperature may also help, but often that is
difficult because of the proximity of other temperature
sensitive detectors. Another standard approach is to add
alcohol, methylal, or water. The oxygen in them is supposed to
slow the growth of the polymer chains. Some people have
suggested adding straight oxygen, but few appear to have done
so. Boyarski[3] has reported success in reversing cathode
aging by adding 200-500 ppm of oxygen. When we finally
added oxygen, it also appeared to be very effective in reversing
aging on the anode.

Another popular approach is to get rid of all hydrocarbons in
the gas. Switching to argon/CO$_2$ or argon/CF$_4$ should eliminate
the type of polymerization that we were seeing. However
argon/CO$_2$ is not as forgiving a chamber gas as argon/ethane
with respect to chamber breakdown.

V. REDUCING AGING IN THE COT

The first step we took was to increase the gas flow into the
chamber from the initial 20 SCFH to 40 SCFH. The higher
flow rate gives a volume exchange every 15 hours.

The next thing we implemented was a gas recirculation
system. The initial goal was to achieve a total flow through the
chamber of 200 SCFH by recirculating 160 SCFH. Cleaning
the recirculated gas was left for the future, but we still expected
to see an improvement because of better gas flow patterns in
the chamber and a dilution of polymers in the inner super-
layers. Gas cleanup was expected to be difficult because of the
large alcohol content that had to be removed from the gas and
then added back into the gas.

Since both the peak and integrated luminosity had significant
increases during this time period, it was difficult to assess how
much these improvements helped. When the input gas rate
was increased from 20 to 40 SCFH, the apparent aging per unit
integrated charge did not appear to change much, but it would
have probably increased if we had done nothing because of the
higher radiation rates. When we added the recirculation, it
appeared to make some improvement, but we did not run long
enough to get a definitive measurement.

However, while installing the recirculation system and
commissioning it, we unintentionally introduced oxygen (from
air leaks) at the 100 ppm level at two well defined points in
time. Eventually we made the right plot (see Fig. 6) to be able
to correlate the oxygen with gain recovery. Adding oxygen
was already on our list of things to try, but we were surprised
by the efficacy with which it worked. Under normal operating
conditions the oxygen level in the COT was $\leq$ 12 ppm; on June

Fig. 5. EDS spectrum of the coating of an aged wire from
the COT. A large carbon and small oxygen peak are seen
along with gold peaks from the wire.
enough air was added to the COT to increase the oxygen level to ~100 ppm. Fig. 7 shows the ratios of super-layer current to the instantaneous luminosity for SL1, SL2, SL4, and SL8 plotted against the integrated charge in SL8 for those times when the inner super-layers were active. The intentional introduction of oxygen started at an integrated charge of 7.0 mC/cm for SL8 (corresponding to ~40 mC/cm for the inner super-layers). Almost two years of gain loss was reversed in less than 10 days while accumulating ~2 mC/cm of charge on the inner super-layers. SL8 appears to have suffered little aging and thus can be used as a reference. The other super-layers show varying degrees of gain loss for much of the run. Fig. 8 plots the average width of the inner super-layers normalized to the average width of SL8 versus the integrated luminosity since the beginning of the run. The dependence on many of the systematic effects such pressure, temperature, and gas mixture cancel in the ratios. This plot shows the same features as Fig. 7.

Probably the oxygen inhibits polymer strings from growing by combining with CH2 radicals or attaching directly to the end of a string\[4\][5]. Aging presumably depends on the balance between the deposition of polymers and the natural ablation that takes place in an avalanche. In this scenario a small amount of oxygen in the gas inhibits polymerization and...
Ablation dominates. However we cannot rule out the possibility that the oxygen is reacting directly with the coating.

VI. STUDIES TO UNDERSTAND THE AGING

Several studies and tests were undertaken to help understand the nature of the aging that we were seeing.

One of the first tests we did was to reverse the gas flow in the chamber. There was concern that there might be contamination from one of the endplates. The aging pattern in z reversed as would be expected if the aging were simply due to polymer buildup from the hydrocarbons in the gas.

Some studies involved looking at the historical data. Based on the HV current data, the gain changed significantly over the first few hours of a store during periods of aging. For this reason, the plots showing the ratio of currents or the ratio of current to luminosity for each store excluded the first 5 hours of the store. There was concern that temperature or charging effects took several hours to stabilize. Fig. 9 effectively shows how the ratio of the gain in the first hour to the gain in the second hour changed for the whole run. During the period of serious aging, the gain was 2-3% lower during the first hour. After the oxygen was added, the gain drop during the first hour decreased to about 0.4%. Some of this small drop could be a temperature effect since the luminosity has been very high since the oxygen was added. The theory is that by the end of a store polymers have built up both on the wires and in the chamber gas. Between stores the polymers in the gas either deposit on the wires or exit the chamber. When the next store begins, the polymer concentration in the gas is lower so that there is reverse aging for the first couple of hours until the polymer concentration rebuilds. This is consistent with our observation that if the chamber HV was kept at its normal operating point after a store was ended, increased gain loss was seen at the beginning of next store.

There were times, often just after a shutdown when the accelerator was not working well, that some gain recovery was observed in the chamber. Fig. 10 shows the ratio of the currents in SL1 and SL2 to the current in an outer super-layer superimposed on a histogram of the charge accumulated per two day period. The period starting at 510 days since the beginning of 2002 is one where the inner super-layers recovered relative to the outer super-layers and also one where the integrated charge per unit time was smaller than usual. There was another period starting at day 444 where there was a small recovery which again was associated with less integrated charge than normal.

Another observation was that the oxygen we added appears to be consumed in the COT during periods of radiation. We add a constant amount of oxygen, but the amount of oxygen observed in the COT exhaust gas decreases by up to 30 ppm depending on the radiation level in the chamber. At this point the variation in oxygen level appears to be larger than we expect and may be due to some artifact of the oxygen sensor. More effort is needed to understand this effect.

Studies were done on some of the aged wires that were removed from the COT in March. They were put in small chambers with a localized source to see if we could reverse age them. One plane of wire was put in a chamber with the

![Fig. 10. Gain in SL1 and SL2 calculated from their HV currents normalized to SL6 versus time in days. The black histogram shows the integrated charge per two days.](image1)

![Fig. 11. Fe$^{55}$ spectra at four different holes in a test chamber constructed with aged wires from the COT. Hole 8 was irradiated with a Sr$^{90}$ in an attempt to recover the gain. The black histogram is the spectra before irradiation, the red at midpoint, and the green at the end.](image2)
standard COT gas mix of argon/ethane (50/50) with 1.7% isopropanol. It was irradiated with a localized Sr\(^{90}\) source and the gain was monitored at several points in the plane by measuring the Fe\(^{55}\) spectrum. It was found the gain made a dramatic recovery underneath the Sr\(^{90}\) source, but it went down elsewhere in the chamber. Fig. 11 shows the Fe\(^{55}\) spectra under hole 8, the irradiated hole, and three other holes away from the source. When the same test was tried with a gas mixture of argon/CO\(_2\), there was again a dramatic improvement under the Sr\(^{90}\) source, but the gain stayed the same or got slightly better elsewhere in the plane. We also put a single aged wire from the COT in a stainless-steel tube, and exposed it to a similar Sr\(^{90}\) source while operating with exhaust gas from the COT that contained about 100 ppm of oxygen. Again we saw a dramatic gain recovery under the source. If we ran these reverse aging tests long enough, the Fe\(^{55}\) spectrum recovered to normal.

After these reverse aging tests, we viewed the wires with a SEM and performed an EDS analysis of the surface. The wires looked clean and the EDS indicated that most if not all of the coating was removed. One point of note is that the EDS analysis of the wire that had been reverse aged in argon/CO\(_2\) showed an anomalously large amount of oxygen.

An attempt was also made to measure the thickness of the coating on the wires. For 40 µm wires we estimate that a 1.0 µm conductive coating should reduce the gain by about 20% and a 2.0 µm coating by 40%. From the SEM pictures of places on the wire where part of the coating has been removed, we got estimates of the coating thickness that vary from 0.3 µm to 2.0 µm. The larger thickness is probably enough to explain the gain loss we saw, but better measurements of the coating thickness are needed to make a statement. Note that if the resistivity of the coating greater than 10\(^6\) Ω-cm, then it may have a significant effect on the gain.

VII. CONCLUSIONS

We found that aging in a clean drift chamber where the primary quenching gas is a hydrocarbon seems to depend on the buildup of polymers in the gas as it flows through the chamber. The deposition of these polymers causes a coating on the wires that reduces the gain. This gain reduction appears to be a multi-step process that does not increase linearly with the integrated charge on the wire. The aging also appears to be temperature dependent. Adding a small amount of oxygen, ~100 ppm in our case, seems to prevent this type of aging and allows such aging to be reversed. This conclusion does not apply to chambers that have non-hydrocarbon contaminants such as silicon.

The oxygen is consumed in a chemical reaction. Traditionally the oxygen in alcohol or water is supposed to inhibit polymerization by combining with CH\(_2\) radicals or attaching to the ends of polymer chains. Apparently oxygen molecules by themselves are much more efficient at doing this. Aging probably depends on the balance between some natural ablation that occurs in avalanches and the deposition of polymers that have built up in the chamber gas. We can’t rule out the possibility that some of the oxygen reacts directly with the wire coating.

These results point out the difficulty of using small test chambers with localized sources to predict aging in larger chambers. They also emphasize the importance of controlling the amount of oxygen in any tests.

VIII. ACKNOWLEDGMENT

The authors would like to thank the Fermilab staff for their vital contributions.

IX. REFERENCES