

## STATUS OF SAMPLE COMBUSTION AND GRAPHITIZATION LINES AT INFN-LABEC, FLORENCE

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**ABSTRACT.** Since installation of the new accelerator at INFN-LABEC in Florence in 2004, the workload has progressively increased in the radiocarbon dating laboratory, requiring a faster sample preparation throughput and greater versatility to allow for a larger variety of treated materials. The “standard” sample preparation line is based on an elemental analyzer (EA) for the combustion of the pretreated samples, and a vacuum line where the CO<sub>2</sub> is collected and then converted to graphite. This line has been recently redesigned while maintaining the EA, since this instrument provides us reliable and fast sample combustion and separation of gases. The volumes were optimized and all the mechanical parts (e.g. the fittings) were changed in order to improve the vacuum level, thus decreasing the possibility of contamination; finally, the number of graphitization reactors was doubled (from 4 to 8). In those rare cases involving samples characterized by a complex nearly graphitic structure (e.g. burnt residues), the EA might not guarantee a complete combustion. For these samples, we successfully tested the new sample preparation line that has been recently installed at LABEC and especially dedicated to aerosol samples. This line is equipped with a custom-made combustion oven able to heat to 1000 °C. The subsequent gas separation is accomplished by chemical and thermal traps. As an example, the dating of organic matter collected from an Etruscan bronze statue will be presented and discussed.

## INTRODUCTION

The INFN-LABEC laboratory in Florence was formally installed in 2004, carrying on the experience of the applied nuclear physics group in the management and the use of small electrostatic accelerators for research studies in the fields of cultural heritage, environmental studies, and material science in general (Giuntini et al. 1995; Chiari et al. 2002, 2005; Massi et al. 2002; Taccetti et al. 2002). Since the beginning, the laboratory has been dedicated to the development of experimental setups for ion beam analysis (IBA) (Giuntini et al. 2007; Lucarelli et al. 2011) and radiocarbon accelerator mass spectrometry (AMS) (Taccetti et al. 2010) measurements, and to many applications in collaboration with archaeologists, art historians, geologists, and environmental scientists.

As far as <sup>14</sup>C is concerned, our laboratory is a small laboratory; in fact, accelerator beam time is shared between AMS and many IBA applications (6 beam lines have been installed so far). Nonetheless, we are able to process a non-negligible number of samples per year, typically about 300. <sup>14</sup>C AMS measurements are performed using the dedicated accelerator beam line (Fedi et al. 2007), whose main features are bouncer sequential injection, stripping with argon at 2.5MV terminal voltage, ion discrimination by magnetic and electrostatic analysis on the high-energy side, and counting of the rare isotopes by a solid state silicon detector. Many materials have been treated so far (see Figure 1), including charcoal, wood, paper, textiles, and bones. Quite recently, applications of <sup>14</sup>C have been also extended to environmental issues, working on shells or foraminifera and on aerosol samples (Fedi et al. 2012; Bernardoni et al. 2013).

This article describes the state of the art of the AMS facilities at LABEC after the first years of activity of the laboratory, focusing in particular on the equipment and the capabilities that the sample preparation laboratory for <sup>14</sup>C measurements has so far acquired. Two different combustion and

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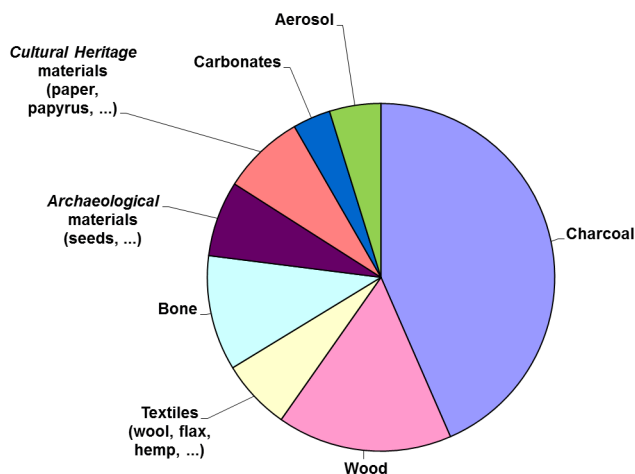


Figure 1 Distribution of the materials of samples treated and measured at INFN-LABEC since installation of the new accelerator (processing of samples like carbonates and atmospheric aerosol has been available for the last 3 yr only).

graphitization lines are installed: one is dedicated to archaeometric and geological applications (i.e. those applications that we can now consider as routinely performed); the other one has been especially designed to be dedicated to aerosol samples. The latter allows us to analyze the different fractions of carbon present in the atmospheric aerosol collected on filters, to perform source apportionment studies thanks to the measurement of  $^{14}\text{C}$  in either the total carbon (TC) or in the separate fractions of organic and elemental carbon (OC and EC, respectively). In the following, we will describe the main characteristics of the 2 combustion and graphitization lines, also discussing possible limitations of their usage. Finally, we will show a case of  $^{14}\text{C}$  dating of organic material collected in an Etruscan bronze statue, showing an example of the flexibility of use of the preparation lines installed in the lab.

### THE COMBUSTION AND GRAPHITIZATION LINE FOR “GENERAL” PURPOSES

The “standard” combustion and graphitization line at INFN-LABEC for general purposes, e.g. archaeometric applications, is principally based on a CN elemental analyzer (Thermo Flash EA 1112) to extract and isolate the gaseous  $\text{CO}_2$  from the sample to be dated, on a vacuum line with a  $\text{LN}_2$  spiral trap to condense the evolved  $\text{CO}_2$ , and on several reactors where the  $\text{CO}_2$  is converted to graphite by the reaction with hydrogen at high temperature (600 °C) and in the presence of iron powder as catalyst (Alfa Aesar iron, pellets <10  $\mu\text{m}$ ). Figure 2 shows the schematic layout of the setup.

The use of the elemental analyzer (EA) to burn samples to be graphitized has some advantages: a fast combustion with the self-built possibility to simply separate  $\text{CO}_2$  from the other evolving products (water, sulfur, nitrogen); the possibility to directly measure the C/N ratio with no need to collect a separate fraction of the sample (this can be indeed useful when preparing bone samples; Sciré Calabrisotto et al., these proceedings); a good reproducibility of the time intervals characterizing the transfer of  $\text{CO}_2$  to the graphitization line, giving us the possibility to easily run the process by an automatic system, too. However, some disadvantages should be recognized as well: some memory effects may occur, which, although kept small by appropriate procedures of “no sample” combus-

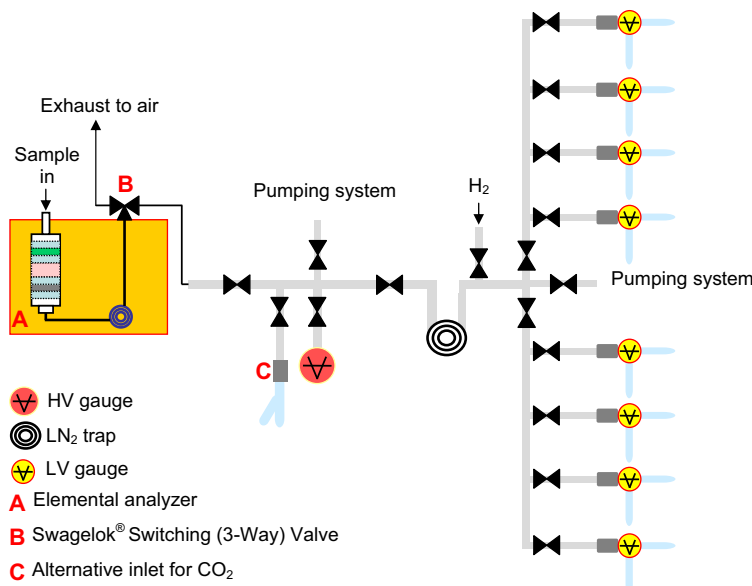


Figure 2 Schematic layout of the combustion and graphitization line for general purposes

tion after each sample combustion to rinse the column. These questions may start to become relevant to the accuracy of the final dating as when burning in sequence very old and modern samples. Moreover, some of the combustion parameters, such as the amount of injected oxygen, can only be regulated within a limited range. Finally, the period of the overall combustion process cannot be set, as it depends on the Dumas combustion characteristics of the EA. This can represent a limit in some cases, e.g. when trying to burn blank carbonate samples as marble or pure graphite-like samples (see below).

This sample preparation line has been recently refurbished and upgraded, to change hardware parts that might have deteriorated since their first installation in 2004 and to enhance the performance (both in terms of quantity of processed samples and reliability of preparation) of the laboratory:

- The EA was coupled to the graphitization line through a new Swagelok® 3-way valve to easily switch the flowing gases; this valve was chosen with a special L-flow path and a reduced size orifice to minimize the possibility of cross-port flow;
- The number of reactors was doubled to graphitize up to 8 samples in parallel;
- A new inlet connection was added to easily transfer the gaseous CO<sub>2</sub> obtained from samples that are not combusted in the EA (see C in Figure 2);
- The first CO<sub>2</sub> spiral trap in the graphitization line was replaced with a flat spiral for a better temperature homogeneity when dipping it into liquid nitrogen (see also Calzolari et al. 2011);
- The use of Viton® O-rings was minimized to improve the vacuum level in the line and to reduce possible effects of contaminations (Swagelok tube fittings with stainless steel ferrules, and weld fittings, were mostly used).

Figure 3 shows the <sup>14</sup>C concentrations measured in IAEA C7 samples (the oxalic acid reference material supplied by the International Atomic Energy Agency with a certified <sup>14</sup>C concentration of  $49.53 \pm 0.12$  pMC) prepared since 2010, when the main changes on the previous setup were applied. The <sup>14</sup>C concentrations reported in the figure were calculated starting from the measured <sup>14</sup>C/<sup>12</sup>C

isotopic ratios, correcting for background and isotopic fractionation ( $^{13}\text{C}/^{12}\text{C}$  ratios also measured along the accelerator beam line), and normalizing to NIST oxalic acid II standard samples. The agreement between the experimental data and the expected value is satisfactory, demonstrating the accuracy of the overall procedure.

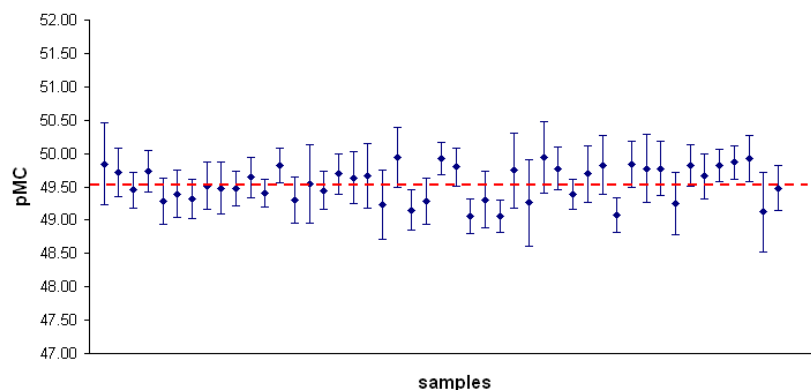


Figure 3 Radiocarbon concentrations measured since 2010 until present (July 2012) in standard IAEA C7 samples, normalizing to NIST oxalic acid II standard samples. The certified  $^{14}\text{C}$  concentration ( $49.53 \pm 0.12$  pMC) is indicated by the dotted line. The experimental uncertainty reported for each sample concentration corresponds to  $1\sigma$ .

Other reference materials are often prepared and measured as secondary standards, and also confirm the accuracy of the measurements:

IAEA C2:  $41.68 \pm 0.43$  pMC (consensus value  $41.14 \pm 0.03$  pMC);

IAEA C5:  $23.03 \pm 0.29$  pMC (consensus value  $23.05 \pm 0.04$  pMC);

VIRI D:  $2860 \pm 46$  yr BP (consensus value  $2836 \pm 4$  yr BP);

VIRI S:  $109.59 \pm 0.82$  pMC (consensus value  $109.96 \pm 0.04$  pMC).

The concentrations and the uncertainties above are reported as average and standard deviations calculated over all the measured samples, respectively. The  $^{14}\text{C}$  concentration measured in processed blank samples is of the order of 0.5 pMC, being limited by the EA rather than the graphitization line.

## THE COMBUSTION AND GRAPHITIZATION LINE FOR AEROSOL SAMPLES

The other sample preparation line installed in the laboratory was specifically designed to meet the requirements of  $^{14}\text{C}$  measurements in aerosol samples. Actually, carbon is one of the major components of aerosol, where it is found as both elemental carbon (EC) and organic compounds (organic carbon, OC). A key point for  $^{14}\text{C}$  measurements in aerosol samples is the possibility to physically separate the different carbonaceous fractions, obtaining from them different samples for  $^{14}\text{C}$  analysis. This type of separation is fundamental in order to get a complete apportionment of the carbonaceous aerosol sources, i.e. to estimate the contributions from fossil fuels and from modern sources and, among the latter, from biomass burning and biogenic sources (Szidat et al. 2006). EC/OC separation is based on thermal protocols including many steps at different temperatures and combustion atmospheres (oxidizing/inert) (Szidat et al., these proceedings).

A detailed description of our aerosol sample preparation line can be found in Calzolari et al. (2011). Briefly, it is based on 4 parts (see Figure 4): 1) cleaning of the carrier gases; 2) combustion of the sample; 3) isolation of the CO<sub>2</sub>; and 4) graphitization. Cleaning of the carrier gases (namely He or O<sub>2</sub> depending on the desired combustion atmosphere) is accomplished by a soda-lime trap after any possible carbonaceous impurity has evolved to CO<sub>2</sub> thanks to a catalyzer. The combustion system guarantees all the features needed for thermal OC/EC separation, as all the combustion parameters (temperature, time, and atmosphere) can be set by the operator and changed according to the adopted separation protocol. The combustion system is based on an oven (specifically designed and in-house built) provided with a catalyzer and a suitable heating coil. With this oven, temperature can be raised up to 1000 °C at a quick rate up to 200 °C/min. Moreover, both the combustion atmosphere and time can be set by the operator. During the combustion, a series of chemical and thermal traps allows the separation of the CO<sub>2</sub> from the other gaseous combustion products and from the carrier gas. Afterwards, the CO<sub>2</sub> trap is filled with He (taking advantage of the shortcut that allows the gases to be injected directly into the trap, bypassing the oven), cooled down to LN<sub>2</sub> temperature, connected to the last part of the line, operating under in-vacuum conditions and devoted to graphitization, and evacuated. Finally, CO<sub>2</sub> is graphitized in a reactor similar to those installed in the sample preparation line for “general” purposes.

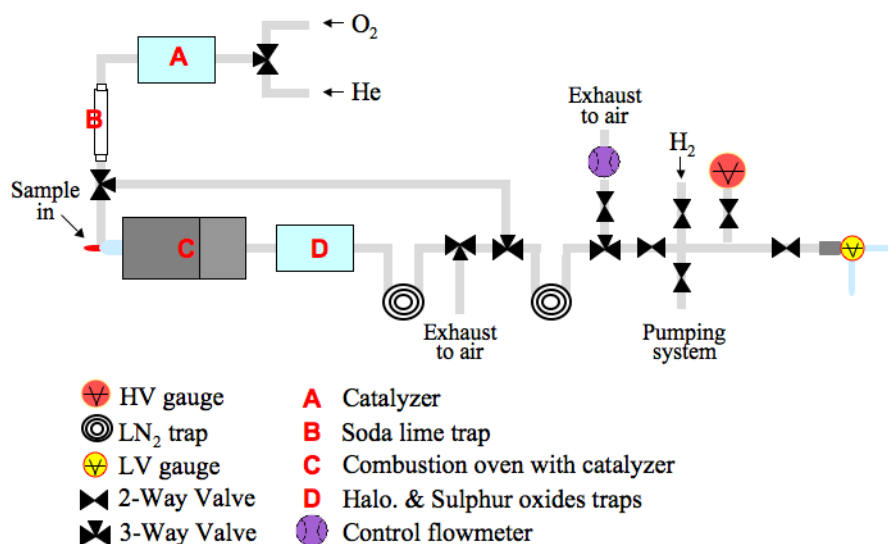


Figure 4 Schematic layout of the sample preparation line dedicated to aerosol samples

Specific combustion protocols were developed for the separation and the analysis of the aerosol carbonaceous subfractions EC and OC (Bernardoni et al. 2013); for the study of the whole carbonaceous fraction (i.e. total carbon, TC), a 20-min combustion at 800 °C, in a 100 cc/min oxygen flow, has been adopted. The sample preparation line was found to have a 100% efficiency for TC samples prepared from different materials (e.g. NIST oxalic acid II, Alfa Aesar graphite, aerosol samples), including in such evaluation the efficiencies for every single step, such as combustion, trapping, transfer, and graphitization (Calzolari et al. 2011).

For 450-μg C samples, the background level was estimated to be about 0.4 pMC (it is worth noting that the expected minimum <sup>14</sup>C content in the aerosol carbonaceous fractions is generally expected to be about 1 order of magnitude higher than this).

### **An Example of Application on Charred Organic Material**

Samples with a complex graphitic structure (e.g. burnt residues) require a highly oxidizing environment, characterized by a large amount of combustive agent and high temperatures for long times, to achieve a complete combustion. According to our experience, for this kind of sample, the flash combustion in the elemental analyzer may not be sufficient. Instead, the aerosol preparation line, in particular the burning oven, although specifically developed for aerosols, allows us to select the combustion parameters with no limitations, leading to a full combustion of the sample.

An example of charred organic material can be shown by some of the organic residues in the clay matrix used to model the body of a bronze sculpture. Many different materials are usually found in the soil, such as animal hair, straw, charcoals, and general charred matter. The latter typically consists of amorphous carbon, and is the result of the combustion at the high temperatures (~400–500 °C) required for realization of the artworks.

To better trace the framework of its origin and its later discovery, characterizing the history of the preservation of this artwork, we were asked to date the organic material collected from the inner part of a famous Etruscan bronze statue, the Chimaera of Arezzo (see Figure 5), now at the National Archaeological Museum in Florence. The statue was discovered near Arezzo (Tuscany, Italy) in 1553, and was immediately taken to Florence at the Medici court. It represents the mythological beast of Chimaera, with the body and the head of a lion, the tail of a snake, and a head of a goat on its back. According to Greek mythology, the Chimaera was a monster, spitting fire from its mouth, which frightened the region of Asia Minor. The statue is known as one of the best examples of Etruscan bronze art and is attributed to a period between the 6th and 5th century BC. However, it is also known that, since its discovery, it has undergone several “restorations” during the centuries. For example, the tail was discovered later and applied to the body only during a restoration in the 18th century AD, even though in the wrong position. Nonetheless, the body of the animal has remained original.



Figure 5 Photograph of the Chimaera of Arezzo, now kept at the National Archaeological Museum in Florence.

For  $^{14}\text{C}$  dating, we collected by scalpel some burnt organic residues from the inner part of the mouth of the statue. First, we observed the collected material under an optical microscope to remove the traces of silicates and metallic compounds, then we treated the organic residues using 1M HCl at high temperature (~80 °C) to remove any carbonates. To reduce the sample to graphite, the aerosol sample preparation line was used, exploiting the TC combustion protocol (see above). We could not

exclude that some metallic residues had been left in the sample matrix and the EA appeared to be unsuitable for the combustion, as discussed above. Primary standard samples (from NIST oxalic acid II), secondary standard samples (from IAEA C7), and blank samples (from Alfa Aesar pure graphite) were also prepared using the aerosol combustion and graphitization line.

Table 1 shows the result of dating. The calibration of the measured  $^{14}\text{C}$  age was performed using OxCal v 4 (Bronk Ramsey 2009) and the IntCal09 calibration curve (Reimer et al. 2009). The result is compatible with the period during which the statue was discovered. This was attributed to the fact that we might have dated not the original organic matrix but something deposited later. This interpretation is consistent with historical documents: in fact, according to the chronicles of the period, the statue was exposed during the celebrations of the Florentine aristocracy, when torches were lighted just inside the Chimaera mouth to reproduce the mythological fire spitting (S Siano, personal communication). We thus expect to have dated those organic residues.

Table 1  $^{14}\text{C}$  dating of the organic residues collected in the inner part of the mouth of the Chimaera of Arezzo. The measured  $^{14}\text{C}$  concentration and corresponding  $^{14}\text{C}$  age are quoted at  $1\sigma$  uncertainty; the calibrated age is quoted at 95% of probability.

	$^{14}\text{C}$ concentration (pMC)	$^{14}\text{C}$ age (yr BP)	Calibrated age
Chimaera	$95.18 \pm 0.52$	$395 \pm 45$	AD 1430–1635

## CONCLUSIONS

Since its start in 2004, AMS activity at LABEC has rapidly increased, and today we are able to process a significant number of samples per year, and to treat most of the materials of interest for  $^{14}\text{C}$  analysis. The sample preparation line for “general” purposes fulfills the requirements for most of the archaeological and geological samples. A recently installed sample preparation line devoted to aerosol analysis allows us to perform  $^{14}\text{C}$  measurements on the separate carbonaceous components of particulate matter. The latter sample preparation line can also find applications for other samples when the choice of the combustion parameters is required, as in the presented case study.

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