



Article X-Ray Dating of Ancient Linen Fabrics

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Abstract: We propose a new method for dating ancient linen threads by inspecting their structural degradation by means of wide-angle X-ray scattering. X-ray dating of a textile sample can be performed nondestructively and on a submillimeter area, e.g., $0.2 \times 0.5 \text{ mm}^2$, exploiting new table-top X-ray micro-sources. A theoretical formula is derived for dating linen samples directly from wide-angle X-ray scattering measurements. Our preliminary results show that X-ray dating results are in agreement with other dating sources, such as the radiocarbon method and historical records, if some conditions are satisfied. Indeed, this new dating method can be applied only to threads not older than about thirty centuries because of the saturation of the structural degradation with age. Moreover, the method can be applied only on textiles in which cellulose degradation is mainly due by natural aging arising from thermal, hydrolytic, photolytic, photochemical, and oxidative processes. Analyses can be repeated several times on the same sample, which remains unaltered for other complementary characterization procedures. The proposed X-ray dating of some ancient linen fabrics opens the way to explore limits and potentialities of this new approach and to further develop a new dating method, alternative to the existing ones for specific applications in archeological studies.

Keywords: radiocarbon dating; X-ray dating; WAXS; cellulose degradation; natural aging; archaeological textiles

1. Introduction

Textiles contain important archaeological information about everyday life, religion, art, technical culture, religion, and history of ancient peoples. Unfortunately, ancient textiles are rare in archaeological sites compared to ceramic, metal, and wood artifacts because of their structural fragility under the action of humidity, temperature, fungi, microbes, and insects. Nevertheless, especially in Egypt or the Near East, there are many cases in which well-preserved textiles have been discovered in crypts and ground caves. Archeological textiles can comprise braided and woven structures from animal or vegetable fibers. Fragments of linen burial cloths indicate that weaving with flax was already used about 8000–9000 years ago [1]. Indeed, flax is a vegetable textile frequently used by populations of the Mediterranean geographical area. One reason is that linen fibers are the most resistant vegetable fibers to environmental aging agents (oxygen, humidity, light, etc.).

Deterioration of vegetable fibers is related to cellulose degradation, the main component of their structure. Cellulose in linen fabrics is a polymer, which contains anhydroglucose units bonded

by β -glycosidic bonds. Aging of textiles can have thermal, hydrolytic, photolytic, photochemical, and enzymatic origins. All these physical and chemical agents degrade cellulose, provoking the scission of the polymeric chains by breaking the β -glycosidic bonds [2–4]. Within any glucose unit, three hydroxyl groups are available for oxidation, leading to a great variety of structural modifications and combinations of the cellulose through oxidative processes. Depending on the particular stress imposed on cellulose, the individual hydroxyl groups in both the polymeric chain and the anhydroglucose units are involved to a varying extent in giving rise to carbonyl and carboxyl groups. These groups are usually introduced in the structure of cellulose through material processing steps, such as bleaching. Furthermore, high energy radiation can cause a considerable increase in carbonyl groups in addition to chain scission, mainly through the action of the generated radical species. For example, the number of carbonyls introduced by γ -irradiation is directly correlated to the applied dosage [5].

Carbonyl and carboxyl group formations in cellulose are also caused by natural aging, which happens at room temperature and humidity. Indeed, the presence of oxygen, often in the presence of light, induces many auto-oxidative reactions. In fact, natural aging of cellulosic fabrics is due to two main processes: hydrolytic cleavage of the glycosidic bond [6–8] and oxidation [9–12], which mainly depends on average room temperature values. Indeed, the role of atmospheric oxygen is crucial. Radical reactions come into play leading to formation of carbonyl or carboxyl groups, to chain cleavage, and thus also to a loss in the degree of polymerization and in the fiber strength.

Recently, new dating methods of textiles have been proposed based on the mechanical analysis such as breaking strength, Young modulus, and loss factor and on opto-chemical analyses (by FT-IR and Raman spectroscopies) as a function of the age of the fabric [13–15]. These new dating methods are possible alternatives to the radiocarbon dating of textiles [16]. Kim et al. [17] investigated the influence of oxidation in the cellulose crystalline domains and showed that the degree of crystallinity of the oxidized cellulose determined by X-ray diffraction decreases according to the oxidation level. This finding, in principle, could allow the obtainment of information about the age of the textile by means of x-ray structural characterization. For this reason, the aim of this paper is to study by means of X-ray diffraction the decrease of the degree of polymerization of cellulose in naturally aged linen fabrics as a function of the aging time, mainly due to thermal and hydrolytic actions which have occurred where they have been preserved for centuries.

In the present paper, by characterizing the textile threads with Wide-Angle X-ray Scattering (WAXS), we have investigated the possibility of extracting structural information suitable for deriving the age of ancient linen fabrics. The work can be summarized in three main parts: (1) the analysis of the WAXS data collected from radiocarbon/historically dated ancient linen threads; (2) the study of the degree of polymerization of cellulose as a function of the aging time; and (3) the X-ray dating obtained by our novel approach compared to other dating of the same investigated linen threads, such as radiocarbon (¹⁴C) or Historical Records (HR). In the discussion of the results, we analyze expected limits of application of the proposed new dating method by verifying the WAXS-derived aging times of linen's threads also from a statistical point of view.

2. Degree of Polymerization as a Function of the Aging Time

Cellulose consists of linear, polymeric chains of cyclic (anhydroglucose) units of which number per chain is called the Degree of Polymerization (*DP*) [3]. During polymerization, the chains of cellulose molecules are organized in both crystalline and amorphous regions, in micro-fibrils, and finally into fibres. The depolymerization, i.e., the increase in the number of Chain Breaks (*CB*), is the main mechanism of cellulose degradation.

The *CB* is proportional to the Scission Fraction of Cellulose Unit (SFCU), which can be related to the *DP* by means of the following equation [3,4]:

$$CB(t) \propto SFCU(t) = \frac{1}{DP(t)} - \frac{1}{DP(t=0)}$$
(1)

where DP(t) is the degree of polymerization at time *t* and DP(t = 0) is the degree of polymerization before the degradation of cellulose starts. *DP* represents the average number of monomers in the cellulose polymer. Therefore, DP(t) gives a measure of the length of cellulose polymer after the aging time *t*. The natural aging process that degrades cellulose in thousands of years can be accelerated in suitable ovens by increasing the temperature to reach the maximum number of *CB* in few thousands of hours. For this reason, by accelerating the process in ovens, the degradation of cellulose is studied under controlled conditions, but there are no data available concerning the *DP* decrease due to thermal, hydrolytic, and oxidative natural aging occurring at room temperature. Threads of ancient fabrics are studied here by WAXS to retrieve the first experimental information about DP(t) or CB(t)for very long natural aging processes.

To relate CB(t) to some quantities measured by WAXS on linen threads, we have defined the fraction of maximum *CB*, a parameter given by the ratio between *CB* at any aging time and the normalized asymptotic value obtained for very long aging times (see Equations (A5) and (A6) in Appendix A):

$$CB_{nor}(t,T_r) = CB(t,T_r)/CB(\infty,T_r) = 1 - \frac{1}{\omega_{DP}^* + (1 - \omega_{DP}^*) \times exp[a_{TDP}(T_o,T_r) \times k_{DP}(T_o) \times t]}$$
(2)

where k_{DP} is the reaction rate at the oven temperature T_o at which the accelerated degradation process has been measured and T_r is the temperature at which fabrics have been preserved during the centuries (i.e., the local average room temperature). Given k_{DP} , T_r , and T_o , Equation (2) depends only on two quantities, the *DP* activation energy (E_{DPa}) and the *DP* maximum fractional loss (ω_{DP}^*). The Arrhenius shift factor $a_{TDP}(T_o, T_r)$ has the following expression [2]:

$$a_{TDP}(T_o, T_r) = exp\left[\frac{E_{DPa}}{R} \times \left(\frac{1}{T_o} - \frac{1}{T_r}\right)\right]$$
(3)

where *R* is the gas constant (see Appendix A for further details). The first task of this work is to obtain information by WAXS about ω_{DP}^* and E_{DPa} for the natural aging of linen, which occurs over many centuries.

3. WAXS Characterization of Ancient Linen Threads

The characterized samples were linen threads, about 1 cm long and 0.2–0.6 mm wide. Some of them are shown in Figure 1. The threads shown in Figure 1 have been taken from fabric samples already studied by other authors [13,15], of which the nomenclature and average ¹⁴C/HR dating are reported in Figure 1 and in the following text. The main data of the characterized samples are summarized in the first two columns of Table 1.



Figure 1. Low-magnification optical microscopy images of some of the analyzed samples under the same illumination conditions: The central sample is of our age (2000 AD). For the ancient samples, in parentheses are shown the average ¹⁴C/HR dating. The analyzed threads are 0.2–0.6 mm wide and a few mm long.

WAXS experiments were performed at the X-ray Micro Imaging Laboratory (XMI-LAB) on all the linen samples reported in Table 1. The laboratory is equipped with an Fr-E+ SuperBright rotating copper anode micro-source (λ = 0.154 nm, 2475 W) coupled through a focusing multilayer optics Confocal Max-Flux (CMF 15–105) to a Small/Wide-angle X-ray scattering SAXS/WAXS (SWAXS) three-pinhole camera equipped for X-ray scanning microscopy. An image plate (IP) detector (250 × 160 mm², with 100 µm effective pixel size) with an offline Rigaku RAXIA-Di reader was employed to collect WAXS data. WAXS data were collected for each linen sample directly mounted on the sample holder with an acquisition time of 1800 s. The spot size at the sample position was about 200 µm. The image plate detector was placed at about 10 cm from the sample, giving access to a range of scattering vector moduli ($q = 4\pi \times \sin \theta/\lambda$) from 1.5 to around 35 nm⁻¹. Two-dimensional (2D) patterns were then converted into one-dimensional (1D) data profiles after the calibration procedure.

Figure 2 displays the 2D WAXS pattern measured on sample D (575 AD) of Figure 1.

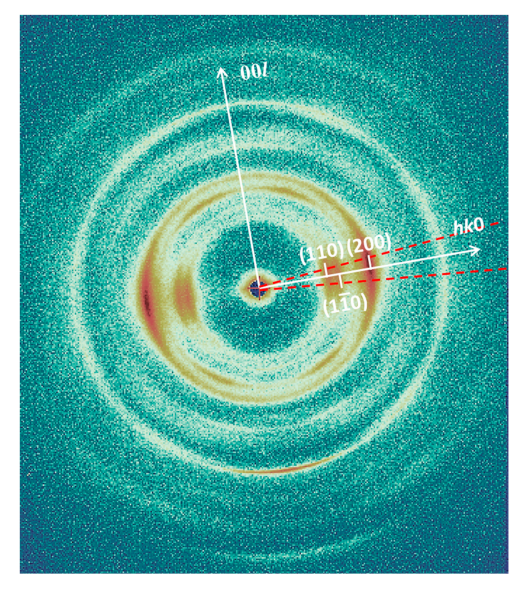


Figure 2. Two-dimensional Wide-Angle X-ray Scattering (WAXS) pattern measured on sample D (575 AD): The main axes of the fiber-diffraction pattern are shown (white lines) together with indexing of main cellulose diffraction peaks.

Figure 3 shows the average scattered intensity of the WAXS patterns of some of the investigated textile samples, integrated in a 10-degree wide angular sector between the two red dashed lines of Figure 2 around the hk0 equatorial fiber-diffraction axis, where h and k are the Miller's crystallographic indexes. This information was extracted from all 2D WAXS patterns of the investigated samples. In particular, in Figure 3, we report the experimental results obtained from sample A (Figure 3, black curve), FII (Figure 3, green light curve), D (Figure 3, red curve), and LII (Figure 3, blue curve), characterized by very different ages. They exhibit clear differences in the scattering profiles, already visible from visual inspection of the curves. The profiles were rescaled to the same integrated intensity in the q-range 1.5–4.0 nm⁻¹, i.e., the region of the incident beam tail. After this rescaling, it is evident that the diffraction WAXS intensity peaks depend on the age of the linen threads. This dependence was used to date the ancient linen threads from WAXS data.

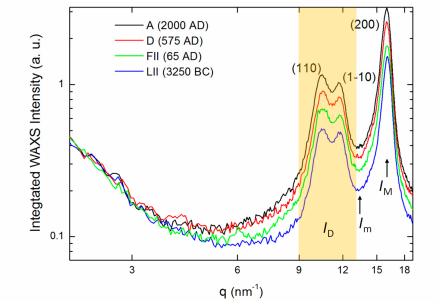


Figure 3. One-dimensional WAXS profiles extracted from the relevant 2D patterns after integration in the angular sector of 10 degrees around the *hk*0 equatorial fiber-diffraction axis in arbitrary units: The cellulose (110) and (1–10) diffraction peaks have been indexed and rescaled to have the same integrated intensity in the q-range 1.5–4.0 nm⁻¹. See the main text for further details and Table 1 for symbols.

Indeed, crystalline cellulose is known to have (1) a diamond-shaped cross section, with exposed (110) and (1–10) crystal faces, and (2) a rectangular-shaped cross section with exposed (200) and (010) faces [18]. Cellulose degradation rates are slightly different for the two crystallographic configurations (diamond- and rectangular-shaped). Moreover, *DP* is not simply related to the crystallinity of cellulose. *DP* could be the same in both crystalline and amorphous regions of cellulose, even if the decrease of the degree of polymerization with age is usually faster in the amorphous regions. However, only spatially ordered chains will contribute to the diffraction peaks. Thus, with WAXS, we can inspect the *DP* of the crystalline regions of the investigated linen threads.

Reaction with cellulosic chains will most likely take place on the surfaces of cellulose crystallites and at defects where water and oxygen can penetrate. Thermal rupture of chains can occur inside crystallites as well as on their outer surfaces based on normal first-order kinetic rupture of glycosidic links. Following rupture of cellulose chains, crystallite volumes will reduce by disordering at surfaces as well as by increasing disorder within crystallites following internal chain thermal rupture. Thus, an average reduction in crystallite volume will ensue but as a consequence of two different mechanisms. The integrated diffraction intensities are proportional to the crystallite volumes and, thus, proportional to the length of the cellulose polymeric chains, i.e., to DP(t) of polymeric chains in the crystalline regions at the aging time *t*. Therefore, the inverse of the integrated intensity under the main cellulose diffraction peaks can be used to obtain experimental information about 1/DP(t). However, from Equation (1), CB(t) is related to 1/DP(t). Thus, by WAXS quantifying the reduction in the intensity of the cellulose diffraction peaks shown in Figure 3, it is possible to obtain experimental information about CB(t).

4. X-Ray Aging Parameter

Table 1 summarizes some experimental quantities obtained by WAXS: the scattered intensity, integrated below the (110) and (1–10) diffraction peaks $\langle I \rangle_{q \in [0.9,1.3]}$ in the scattering vector q-range = 9.0–13.0 nm⁻¹ (yellow region of Figure 3); the maximum intensity in the entire q-range, dominated by the intensity of the (200) peak (*I*_M); and the minimum background intensity between the (1–10) and (200) diffraction peaks (*I*_m). The symbol <...> stands for the average of the scattered intensities in the indicated q-range. The term $\langle I \rangle_{q \in [0.9,1.3]}$ stands for the intensity in an angular sector of 10 degrees around the hk0 axis averaged in the scattering q-range from 0.9 to 1.3 nm⁻¹ and is referred in the following as *I*_D.

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Sample Name	Sample Description of Fabric (Provenience)	Year of Manufacture Based on ¹⁴ C Dating or Historical Records	Scattered Intensity under the (110)–(1– 10) Peaks	Maximum Main Diffraction Peak Intensity	Minimum Main Diffraction Peak Intensity	Main Diffraction Peak: Max-to-Min Ratio	Aging Factor
			ID <i>q∈ [9.0,13.0]</i>	Iм I(200)	Im Iq=13.3	RMm IM/Im	$\frac{AF}{\frac{I_M}{I_m}} \times \frac{1}{I_D}$
В	New, raw flax	2000 AD	0.948 ± 0.009	3.63 ± 0.01	0.507 ± 0.002	7.16 ± 0.03	7.56 ± 0.08
А	New, bleached	2000 AD	0.996 ± 0.008	4.186 ± 0.009	0.517 ± 0.005	8.1 ± 0.1	8.1 ± 0.1
DII	Medieval, ¹⁴ C dating, Jericho (Israel)	997–1147 AD	0.732 ± 0.009	2.941 ± 0.007	0.471 ± 0.004	6.24 ± 0.04	8.5 ± 0.1
D	Medieval, ¹⁴ C dating, Fayyum (Egypt)	544–605 AD	0.833 ± 0.008	3.57 ± 0.01	0.460 ± 0.002	7.76 ± 0.04	9.3 ± 0.1
FII	From a mummy, HR dating (Siege of Masada), (Israel)	55–74 AD	0.641 ± 0.008	2.41 ± 0.01	0.366 ± 0.004	6.58 ± 0.07	10.3 ± 0.2
NII	From a mummy, ¹⁴ C dating, Engedi (Israel)	350–230 BC	0.538 ± 0.007	1.91 ± 0.01	0.308 ± 0.004	6.20 ± 0.09	11.5 ± 0.2
Е	From a mummy, ¹⁴ C dating, Egypt	405–345 BC	0.416 ± 0.005	1.799 ± 0.007	0.380 ± 0.004	4.7 ± 0.1	11.4 ± 0.2
HII	From a mummy, ¹⁴ C dating, Thebes (Egypt)	1000–720 BC	0.825 ± 0.007	3.90 ± 0.01	0.41 ± 0.005	9.5 ± 0.1	11.5 ± 0.2
LII	From a mummy, ¹⁴ C dating, Egypt	3500–3000 BC	0.649 ± 0.007	2.76 ± 0.01	0.367 ± 0.002	7.52 ± 0.05	11.6 ± 0.1

Table 1. Data obtained from the analysis of the 1D WAXS profiles: Sample description and age are obtained from the literature [13,15]. The nomenclature of the samples is the same used in the cited papers. AD = Anno Domini; BC = Before Christ. Scattering vector q-values are given in nm⁻¹.

From these experimental quantities, we extracted the Aging Factor (AF), which is proportional to the inverse of I_D , i.e., proportional to the inverse of DP(t). The integrated intensity under every diffraction peak is also proportional to the number of crystalline cellulose chains, which diffract photons in well-defined scattering directions, defined by the Bragg law. Therefore, in the definition of AF, it is necessary to insert a quantity proportional to the number of cellulose chains, which cause diffraction. In turn, this quantity depends on the actual thickness of the sample, which changes for every sample and comprises a different number of fibres, as shown in Figure 1. This contribution to the intensity of the peaks can be taken into account by the maximum-to-minimum ratio of the main (200) peak, evaluated as follows:

$$R_{Mm} = \frac{I_M}{I_m} \tag{4}$$

where I_M is the maximum intensity corresponding to the (200) diffraction peak and I_m is the minimum scattered background intensity that occurs at about q = 13.3 nm⁻¹ between the (1–10) and (200) peaks.

Thus, the Aging Factor (AF) is defined by the following equation:

$$AF = R_{Mm} \times \frac{1}{I_D} \tag{5}$$

which is a quantity proportional to the actual thickness of the sample and to the inverse of DP(t). We have considered only natural aging to determine unknown quantities such as ω_{DP}^* and E_{DPa} , from which the cellulose degradation process depends on. Natural aging happens when the textiles have suffered only thermal, hydrolytic, photochemical, and oxidative degradation processes at stationary room conditions over the centuries.

Our experimental tests indicate that samples in which cellulose has been degraded by natural aging (thermal, hydrolytic, oxidative, and photochemical processes at room temperature and humidity) are characterized by *R*_{Mm} values larger than 4. Let us note that the maximum-to-minimum ratio of the main (200) peak *R*_{Mm} can be also indirectly related to the degree of crystallinity of the cellulose through the following relation [19]:

$$C = \frac{I_M - I_m}{I_M} = 1 - \frac{1}{R_{Mm}}.$$
 (6)

Although the above relation usually overestimates the degree of crystallinity of cellulose [20], it indicates that, below a given level of crystallinity, corresponding to R_{Mm} values smaller than 4, causes other than natural aging have probably contributed to the degradation of cellulose. Indeed, the *DP* maximum fractional loss and the *DP* activation energy values change under the action of enzymes or acid/basic solutions, which strongly accelerated cellulose degradation. Our tests indicate that even an empirical and approximated evaluation of *C*, through Equation (6), allows a rapid comparison of cellulose samples [20] to make a preliminary judgement if the linen fabric has suffered mainly a natural aging process and so to discard samples that could have suffered more aggressive degradation processes.

The intervals reported in columns 4–7 of Table 1 by means of the " \pm " symbol are uncertainty intervals related to the repeatability of the WAXS measurement in the same point of the same sample. The interval values reported in columns 7–8 of Table 1 have been obtained by propagating the corresponding uncertainties. They correspond to a single measurement of 1800 s on a single point of the sample. For homogeneous degradation due to natural aging of the sample along its extension, it is possible to derive *AF* values which do not depend on the particular position on the thread within the uncertainty range of the measurements. Indeed, either repeating the measurement on the same point or changing the point of analysis along the thread, the same results are obtained, within the experimental errors. Therefore, if the sample degradation is homogeneous along all its extension, the averaging of many measurements allows for the reduction of the uncertainty intervals obtained relative to a single one. For the representativeness of the WAXS characterization of a single

thread for the whole fabric, see the Discussion section. We checked that repeating the WAXS characterization, in the same point of the linen threads after an irradiation of 15 hours with the X-ray beam (8 KeV of energy) used in the measurements does not cause a measurable structural degradation within the experimental errors. Moreover, the average of four measurements of 1800 s, which halves the corresponding uncertainties, is already sufficient to reduce the uncertainty interval enough to become comparable with other source of errors, like the uncertainty on the average room temperature.

AF values obtained by a single measurement of 1800 s are reported in the last column of Table 1, and plotted in Figure 4. From Figure 4, the almost linear growth of the parameter *AF* as a function of the aging time estimated by ¹⁴C/HR dating up to about 25–30 centuries, after which saturation occurs, is evident. This finding can be related to the evolution equation of *CB*, reported in Appendix A [3–4], as a function of the aging time *t*.

From Equation (1), it follows that CB(t) is proportional both to DP(t), the degree of polymerization at time t, and to DP(t = 0), the degree of polymerization before the degradation of cellulose. Therefore, we can introduce an aging parameter related to $AF-AF_{min}$, a quantity that is proportional to 1/DP(t)-1/DP(t = 0), where AF_{min} is the minimum value of the aging factor, corresponding to the maximum value of DP, that is DP(t = 0), without any degradation due to natural aging. This value could be obtained by the contemporary linen sample of raw flax, for which we can put the aging time t = 0 on the scale of centuries. Therefore, $AF-AF_{min}$ is proportional just to CB(t). Moreover, we have introduced a normalized CB_{nor} in Equation (2) to have an a-dimensional quantity to evaluate natural aging of cellulose.

For the same aim, we can introduce a normalized Aging Parameter (*AP*) obtained from WAXS measured values, as follows:

$$AP = \frac{AF - AF_{min}}{AF_{max} - AF_{min}} \tag{7}$$

where the AF_{\min} is the minimum value corresponding to contemporary raw flax and AF_{\max} is the maximum value obtained for very long aging times.

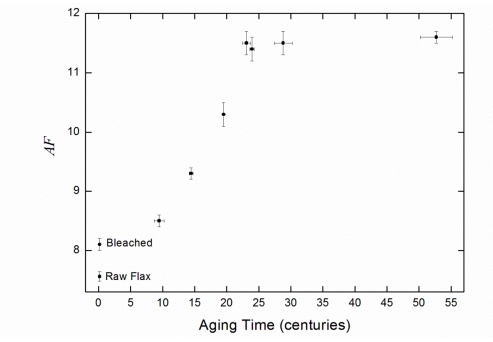


Figure 4. Aging Factor (*AF*) values as a function of the ¹⁴C/HR dating of the investigated linen threads: Error bars are four standard deviations (-2σ , 2σ).

Given the proportionality of $AF - AF_{min}$ to 1/DP(t)-1/DP(t = 0), we can consider AP as an experimental evaluation of CB_{nor} obtained by WAXS. Thus, the evolution equation of CB_{nor} can be used to fit the aging parameter AP values derived from AF values determined by the WAXS measurements (see Figure 4) as a function of the aging time t. In this way, we can experimentally determine the DP maximum fractional loss , ω_{DP}^{*} , and the DP activation energy, E_{DPa} , for the natural aging of linen that occurred during very long periods of time at about stationary conditions over a time scale of many centuries. The evaluation of these quantities is a prerequisite to perform an X-ray dating of ancient linen threads. Whenever the fraction of maximum CB, i.e., the CB_{nor} values, can be quantitatively derived by AP, the X-ray aging times obtained can be compared with those obtainable by other dating methods.

5. X-Ray Determination of Linen Aging Parameters

The normalized *AP* values, obtained by applying Equation (7) to the mean of the measured *AF* values, are reported in Table 2, together with the age of the samples determined by ¹⁴C/HR dating. Sample A has been discarded by the analysis because bleaching processes have introduced a nonnatural aging, i.e., an accelerated *DP* reduction due to the chemical action of oxygen used for the decolorization of the textile [2]. This finding is evident also by our WAXS characterization, which gives an *AP* value of the bleached thread (sample A) larger than that obtained by the raw flax sample (sample B). Indeed, the oxidative bleaching of the fiber has introduced a number of *CB*, causing a certain level of structural degradation of the fabric. For further considerations about the action of acid, basic, enzymatic chemical processes on the structural degrade of cellulose, see the Discussion Section.

For $T_o = 273.15 + 90$ K, the reaction rate k_{DP} of the linen DP degradation, usually given in hours⁻¹ (denoted in the following as h^{-1}), is reported to be $k_{DP}(273.15 + 90) = 0.0022 h^{-1}$ [4]. Natural aging depends on the average room temperature at which cellulose degradation occurs. To evaluate the temperature excursions for longer time periods, we can rely on a detailed analysis of tree rings' growth [21]. A 100-year spline filter shows that the global mean temperature oscillations in the last two millennia have been contained in a range of ±0.9 °C. The duration of these warmer or cooler periods of large temperature variations has been at most about two centuries. Moreover, a long-term cooling trend of about 0.6 °C has been manifested in the last two millennia, from the beginning of the Christian Era until the beginning of 1900 [21]. The average room temperature of the last century is very similar to that of the Roman period about 2000 years ago. The average room temperature in the last century is available today for many countries [22]. For Egypt, West Bank, and Israel the average temperatures in the last century have been 22.5 °C, 19.4 °C, and 19.5 °C, respectively. Therefore, due to the results reported in weather data banks [22], these values can be considered the average room temperatures of the last two millennia within small variations, indeed, with a reduction of about 0.6 °C in the last two millennia, divided between twenty centuries, leading to 0.03 °C variations per century. The variations of ±0.9 °C with a maximum duration of about two centuries averaged over a longer period of two millennia, as it was applied a 1000-year spline filter to climate data, are equivalent to a fluctuation of about 0.09 °C per century. Therefore, within an indetermination of about $\Delta T_r = 0.1$, the temperature of the last century can be considered representative to the last two millennia.

Table 2. Age of the samples derived by the ¹⁴C/HR dating, normalized aging parameter AP values obtained by the WAXS measurements, and average room temperature (°C) at which the samples have been degraded.

Sample Name	Aging Time (centuries)	Aging Parameter $AP \pm 2\sigma_{AP}$	Average Room Temperature <i>Tr</i> (°C)
В	0.2 ± 0.1	0.00 ± 0.01	/
DII	9.5 ± 0.8	0.23 ± 0.01	21.5
D	14.4 ± 0.3	0.43 ± 0.01	21.0

FII	19.55 ± 0.1	0.68 ± 0.02	21.0
NII	23.1 ± 0.6	0.98 ± 0.025	23.5
Е	23.9 ± 0.4	0.95 ± 0.025	22.5
HII	28.8 ± 1.4	0.98 ± 0.025	22.5
LII	52.7 ± 2.5	1.00 ± 0.01	22.5

However, local microclimate conditions can cause differences in the average room temperatures of some degrees with respect to average values of the surrounding regions. This happens in the Dead Sea Valley, the deepest depression of the Earth. Two samples—DII (Jericho) and NII (Engedi)—come from this depression. Mean annual temperatures for these localities in a temporal range of some decades are available. For Jericho, we found 21.5 °C [23]. For Engedi, we found a mean annual temperature of 23.9 °C for the years 1982–2012 [24], which should imply about 23.5 °C for the average room temperature in the last century due to the increasing average room temperatures of the last few decades. For Fayyum and Masada, we found about 21.0 °C. In the last column of Table 2, we reported temperatures expressed with an accuracy of 0.5 °C. They can be considered the average values at which the fabrics were aged during the time. Room temperatures in Table 2 have been given in °C, a unit commonly used for meteorological and climate studies. Conversion in Kelvin allows their use in the theoretical formulae of cellulose degradation.

There are no available data about the relative humidity during the last two millennia for all the localities where the considered linen fabrics aged. Nevertheless, the studied samples come from regions with similar climate, semidesert, with an average relative humidity ranging between 50 and 60%. Averaging over many centuries leads to very similar relative humidity values. k_{DP} (273.15 + 90) values between 0.0021 h^{-1} and 0.0022 h^{-1} have been measured for cotton and linen at 90 °C and 50% of relative humidity [4]. An increment of 10% of the relative humidity, from 50% to 60%, causes an increment of about 10% of the cellulose k_{DP} [25]. Therefore, we can put k_{DP} (273.15 + 90) \cong 0.0023 \pm 0.0001 h^{-1} both for cotton and linen at a relative humidity ranging between 50% and 60%, with an error on the k_{DP} value related to an error of about 5% on an average relative humidity value of about 55 \pm 5%.

From Equations (A5) and (A6) reported in Appendix A, since AP values can be considered an experimental evaluation of CB_{nor} obtained by WAXS, a relation between the aging time t and the other quantities involved in the natural degradation process of the cellulose can be obtained:

$$t = \frac{1}{k_{DP}(T_o)} \times ln \left[\frac{1 - \omega_{DP}^* \times (1 - AP)}{(1 - \omega_{DP}^*) \times (1 - AP)} \right] \times exp \left[-\frac{E_{DPa}}{R} \times \left(\frac{1}{T_o} - \frac{1}{T_r} \right) \right]$$
(8)

Through Equation (8), an evaluation of the natural aging time of fabrics' threads can be calculated directly from the measured *AP* values and the estimated average room temperatures T_r , given the measured $k_{DP}(T_o)$ at $T_o = 90 + 273.15 K$ and estimated the values of E_{DPa} and ω_{DP}^* .

To estimate E_{DPa} and ω_{DP}^* , we use the experimental data of sample FII, aged at an average room temperature of about 21.0 °C (see Table 2). Its dating has been derived by HR (siege of Masada). The *AP* values, reported in Table 2, have been determined by the WAXS *AF* values shown in Figure 4. Let us note the linear dependence of *AF* and, thus, of *AP*, if *t* ranges from 10 to about 20 centuries. Indeed, if we consider also the sample just before of FII, sample D, in particular, if we consider the aging time's region corresponding to the two samples D and FII, we note a linear growth of the *AF* values (see Figure 4). Since *AP* values are proportional to *AF* values and *AP* values can be considered an experimental evaluation of *CB*^{nor}, the linear growth of *AF* values in the aging time's region of the D and FII samples will be related to the slope of *CB*^{nor}(*t*, *T_r*), at half of its maximum value. In Appendix A, we have derived a relation which relates the *DP* activation energy E_{DPa} to this slope (Equation (A10)):

$$E_{DPa} = \frac{RT_oT_r}{T_o - T_r} \times ln \left[\frac{4 \times \frac{\Delta CB_{nor}(t, T_r)}{\Delta t}}{k_{DP}(T_o)} \right]$$
(9)

From samples D and FII, we can estimate this slope:

$$\frac{\Delta CB_{nor}(t,T_r)}{\Delta t} = \frac{\Delta AP}{\Delta t} \cong \frac{0.68 - 0.43}{19.55 - 14.4} \cong 0.049 \pm 0.003 \tag{10}$$

From Equations (9) and (10), we obtain an estimated *DP* activation energy E_{DPa} of 118.9 ± 0.8 kJ/mole, a value that falls in the range of activation energies measured for the degradation of cellulosic materials [25,26]. We can determine also $\omega_{DP}^* = 0.953 \pm 0.001$ by imposing that Equation (8) leads to 19.55 ± 0.1 centuries as the aging time for the sample FII, of which the dating has been obtained by HR. Our approach has allowed, for the first time to our knowledge, the determination of the factors which affect the natural aging of linen cellulose at room conditions of temperature and humidity, a process which happens very slowly, requiring period of times of many centuries or thousands of years.

6. X-Ray Dating

In the previous section, in order to obtain an estimate of the values of a_{DP}^* and E_{DPa} , we have used the HR dating of sample FII together with the evaluation of the slope of $CB_{nor}(t, T_r)$ at half of its maximum value, obtained by using the ¹⁴C dating of sample D. ¹⁴C dating values of the other samples DII, NII, E, and HII have not been used. Therefore, they are useful for a reliability check of X-ray dating obtained by Equation (8), based on the WAXS experimental evaluation of CB_{nor} given by *AP*. Sample LII is not useful for the check of reliability of X-ray dating because it is too ancient and has reached the maximum *AP* value. In addition, the contemporary sample B is not useful for the check of reliability of X-ray dating because it has not suffered any aging degradation process. In fact, because *DP* degradation has an asymptotic behavior as a function of *t*, we can associate a finite aging time value only up to a maximum *AP* value of 0.99 because, for *AP* \rightarrow 1, Equation (8) is indeterminate. For this reason, for sample LII, we cannot obtain any X-ray dating, and for *AP* values very close to 1, the uncertainty interval of the determined aging time is expected to be very large.

The results obtained by Equation (8) and by the measured *AP* values for $k_{DP}(T_o) = 0.0023 \pm 0.0001 h^{-1}$, $\omega_{DP}^* = 0.953 \pm 0.001$, and $E_{DPa} = 118.9 \pm 0.8$ kJ/mole have been summarized in Table 3. The time intervals reported in the last column of Table 3 should correspond to a 95% level of confidence. The larger 95% right confidence levels of samples NII and HII with respect to the left-hand values are due to the saturation of the *CB*_{nor} curve towards 1 for large aging times. Indeed, these two samples are both characterized by very large *AP* values of about 0.98, and the width of their +2 σ uncertainty interval (right-hand value) is about 70% larger than the -2 σ interval (left-hand value). Further statistical insight about this point will be discussed in Appendix B.

The comparison with the dating of the fabrics obtained by ¹⁴C confirms that the X-ray dating reported in Table 3 could be a valid and reliable new method to estimate the time when ancient fabrics, which have suffered natural aging, were made because any information about the dating of samples DII, NII, E, and HII has not been used to perform the corresponding X-ray dating. In fact, the obtained X-ray dating can be considered conditioned almost totally by the HR dating of sample FII and almost completely independent by the ¹⁴C dating of the other samples. Indeed, the estimate of the slope of the *CB_{nor}* curve, i.e., of the fraction of maximum *CB*, at half of its maximum is partially dependent only on the ¹⁴C dating of sample D. In the third column of Table 3, for X-ray aging times, we show in parentheses the $[-2\sigma, 2\sigma]$ interval which, for older samples, is not symmetric with respect to the median value due to saturation effects in the decreasing degree of polymerization of cellulose, which seems to stop after several thousands of years. For details about the ¹⁴C date ranges reported in the third column of Table 1, one can refers to previous works [13,15]. To facilitate the comparison with the X-ray dating results, the ¹⁴C dating values reported in the second column of Table 3 have

been obtained by expressing the data ranges of the third column of Table 1 as a median value with a $[-2\sigma,2\sigma]$ interval. However, it should be noted that, due to the non-monotonic nature of the ¹⁴C calibration curve, ¹⁴C dating very often leads to the impossibility to summarize the data range interval as a median value with symmetric left and right 95% confidence level limits, since the calculated age distribution can be also bi- or trimodal. Therefore, the ¹⁴C aging time median values reported in Table 3 have to be considered only as equivalent values to allow a direct comparison with the X-ray corresponding dating values.

Sample Name	¹⁴ C/HR Aging Time (95% Confidence Level; 2 Standard Deviations Each Side)	X-Ray Aging Time Obtained by Equation (8) (in Parentheses: Left and Right 95% Confidence Levels; 2 Standard Deviations Each Side)
DII	9.5 ± 0.8	9.4 [-0.7,0.7]
D	14.4 ± 0.3	14.5 [-1.0,1.0]
FII	19.55 ± 0.1	19.6 [-1.5,1.5]
NII	23.1 ± 0.6	23.5 [-2.9,5.1]
Е	23.9 ± 0.4	23.9 [-2.3,2.8]
HII	28.8 ± 1.4	27.7 [-3.6,6.0]

Table 3. Comparison of ¹⁴C/HR dating and X-ray dating obtained by Equation (8).

Figure 5 summarizes the results reported in Table 3 and shows the effect of the average room temperature on the increment of *CB* in the natural aging of linen threads. In Appendix B, the relationships between ¹⁴C and X-ray dating have been further analysed by a suitable statistical analysis of data.

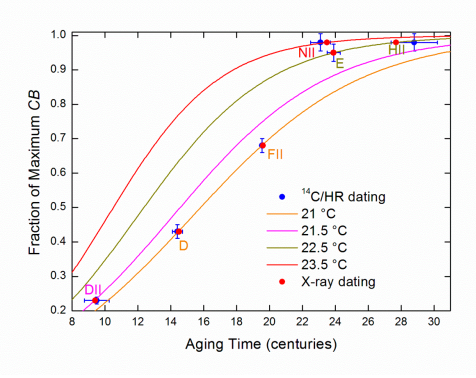


Figure 5. Comparison between ¹⁴C/HR and X-ray dating: *CB_{nor}* curves corresponding to the different average room temperatures of the samples have been plotted with different colors. Error bars (blue) correspond to the ¹⁴C/HR dating (blue circles). Error Bars of X-ray dating (red circles) have not been shown for clarity.

7. Discussion

Our analysis has shown that ancient linen threads that have suffered natural aging due to the oxidative, photochemical, and hydrolytic actions at room temperature and humidity over very long period of times of many centuries can be dated by WAXS characterization of the structural degradation of the cellulose fibers. The X-ray dating results are in agreement with the ¹⁴C dating. Additionally, the measurements are not destructive. These findings allow comparison of different dating methods on the same piece of textile, if needed.

The method does not need any sample preparation, requires just a few hours of measurements, and costs a few hundreds of euros to realize the whole X-ray characterization and data analysis of a single sample. Exploiting table-top X-ray micro-sources, dating of a textile sample can be performed also on a submillimeter area, e.g., 0.2×0.5 mm². However, this possibility is based on the assumption that the ancient textile fabric is homogeneous in its physical/chemical/structural characteristics, i.e., the analyzed small thread is representative of the full textile. This important assumption is critical for other dating methods too. For example, in some cases, ¹⁴C dating of textile fabrics suffers from the impossibility to eliminate all textile carbon contaminants through the standard procedure of sample cleaning, thus leading to possible wrong dating [16]. Therefore, the possibility to perform dating of a textile by using complementary sources of information could be very useful to solve eventual doubtful cases.

Our WAXS measurements indicate that even an approximated but rapid evaluation of the degree of crystallinity of the analyzed sample can allow the selection of ancient threads that most probably have suffered only natural aging over the centuries. These threads are characterized by a higher crystallinity than through other causes of aging (enzymatic and acid/alkaline agents) which accelerate the degradation of the textile with respect to natural aging at room temperature and humidity.

The proposed approach applied to linen threads can be generalized to any textile fabrics. The analyzed threads should be made by the same kind of natural fibers. Indeed, fabrics and paper made by other textile fibers (cotton, hemp, jute, kozo, etc.) could be characterized by *DP* parameters and activation energy values that can be different from linen. Therefore, if one considers a textile different from linen to predict its age by means of the theoretical formula given by Equation (8) and a WAXS characterization of its threads, the following quantities have to be either experimentally determined or known: $k_{DP}(T_o)$; ω_{DP}^* ; and E_{DPa} . However, cotton and linen seem to be characterized by very similar reaction rates, measured with accelerated aging in oven experiments. Therefore, when applying the new dating method to ancient cotton threads, only the *DP* fractional loss and the *DP* activation energy involved in natural aging of the textile, another ancient sample of the same textile of which dating is known from other reliable sources (e.g., ¹⁴C/HR dating) should be available.

In summary, our results have shown that the independent X-ray dating of four ancient linen threads is in agreement with the corresponding ¹⁴C dating. This result shows that ¹⁴C and X-ray methods, if necessary, could be used together to increase the reliability of dating measurements in order to determine the possible presence of either inhomogeneity or impurity in the sample, which could affect the dating resulting by a specific method. The agreement obtained among ¹⁴C and X-ray dating has been also quantitatively analysed by a novel statistical analysis of the data (see Appendix B), showing that the two methods of estimating the aging time are compatible within the experimental uncertainties for the linen samples analyzed here.

8. Conclusions

We propose a novel method for X-ray dating of textiles in which cellulose degradation is due to natural aging, i.e., to thermal, oxidative, hydrolytic, and photochemical processes at room temperature and humidity conditions. In particular, with this approach, we have experimentally determined by means of a WAXS characterization of some threads the *DP* maximum fractional loss and the *DP* activation energy for the natural aging of linen. Natural aging is a degradation process of

cellulose, which very slowly reduces the degree of polymerization, *DP*, requiring many centuries or thousands of years to reach saturation.

The method is not destructive, giving the possibility to analyze the same sample with other experimental techniques. The preliminary results discussed in the paper show that the proposed X-ray approach could be considered to yield novel, independent, and complementary information available to integrate and compare with already existing dating methods on ancient textile fabrics. Further studies will help to clarify the potentiality of the new X-ray dating method proposed here. In particular, the effects of mordant, dyes, and ancient beating procedures of extraction of the textile fibers could, in principle, affect the initial values or the time evolution of the degree of polymerization of cellulose. Specific studies on these topics would allow the clarification of the limits of applicability of this new proposed dating method of ancient textile fabrics.

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Appendix A: Evolution Equation of Chain Breaks in Cellulose

The *DP* fractional loss can be estimated as:

$$\omega_{DP} \equiv 1 - \frac{DP(t)}{DP(0)} = \omega_{DP}^* \times (1 - e^{-k_{DP}t})$$
(A1)

where ω_{DP} is the accumulated *DP* loss at time *t* compared to the initial value *DP*(0); ω_{DP}^* is the *DP* maximum fractional loss; and k_{DP} is the reaction rate of the *DP* degradation. Measures of *DP* loss for linen as a function of time are reported at 90 °C temperature and 50% relative humidity [3]: k_{DP} (90 °C = 363.15 K) = 0.0022 h⁻¹.

The Chain Breaks (*CB*) for linen, in mmol/100 g of cellulose, can be estimated by the *DP* degradation [4]:

$$CB(t) = 1235 \times \left(\frac{1}{DP(t)} - \frac{1}{DP(0)}\right) \tag{A2}$$

From Equations (A1) and (A2), we get the following:

$$CB(t) = \frac{1235}{DP(0)} \times \frac{\omega_{DP}^* \times (1 - e^{-k_{DP}t})}{1 - \omega_{DP}^* \times (1 - e^{-k_{DP}t})}$$
(A3)

We define a normalized *CB* value with respect to its maximum value obtained for very long aging times: $CB_{nor}(t) = CB(t)/CB(\infty)$. After some straightforward calculations, from Equation (A3), the normalized *CB* value can be written as follows:

$$CB_{nor}(t,T) = 1 - \frac{1}{\omega_{DP}^* + (1 - \omega_{DP}^*) \times e^{k_{DP}(T)t}}$$
(A4)

In Equation (A4), the dependence on the temperature *T* has been reported explicitly. For $t \to 0$, one obtains $CB_{nor}(0) = 0$. For $t \to \infty$, one obtains $CB_{nor}(\infty) = 1$.

The effect of the high oven temperature used to measure k_{DP} (90 °C = 363.15 K) [4] is to accelerate the degradation of cellulose with respect to the natural aging at room temperature. The k_{DP} value at room temperature is related to the reaction rate of the *DP* degradation at 90 °C by an Arrhenius shift factor $a_{TDP}(T_o, T_p)$ [2], depending on the activation energy [3]:

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$$k_{DP}(T_r) = a_{TDP}(T_o, T_r) \times k_{DP}(T_o) = exp\left[\frac{E_{DPa}}{R} \times \left(\frac{1}{T_o} - \frac{1}{T_r}\right)\right] \times k_{DP}(T_o)$$
(A5)

where E_{DPa} is the *DP* activation energy; T_o is oven absolute temperature [4]; T_r is average absolute room temperature during the centuries at which the linen fabrics have been aged; and *R* is the gas constant (8.314 J/mol/K).

By inserting Equation (A5) into Equation (A4), we finally obtain an equation to fit the normalized *AP* experimental values reported in the Results section:

$$CB_{nor}(t,T_r) = 1 - \frac{1}{\omega_{DP}^* + [1 - \omega_{DP}^*] \times exp[a_{TDP}(T_o,T_r) \times k_{DP}(T_o) \times t]}$$
(A6)

Equation (A6) is dependent only on two free parameters, E_{DPa} and ω_{DP}^* , once the average room temperature T_r has been decided.

From the time derivative of Equation (A6), we obtain the following:

$$\frac{\Delta CB_{nor}(t,T_r)}{\Delta t} = \left(\frac{1}{1 - CB_{nor}} - \omega_{DP}^*\right) \times (1 - CB_{nor})^2 \times k_{DP}(T_r)$$
(A7)

When $\triangle CB_{nor}$ is close to 0.5, from Equation (A7), we obtain

$$\frac{\Delta CB_{nor}(t,T_r)}{\Delta t} = (2 - \omega_{DP}^*) \times \frac{k_{DP}(T_r)}{4}$$
(A8)

Now, let us consider that, for natural aging of thousands of years, ω_{DP}^* is quite close to one. Indeed, after 4000 hours in an oven at 90 °C, the measured *DP* values for linen decrease from an initial value of 9600 down to 900 [3], implying that, for natural aging of thousands of years, ω_{DP}^* should be greater than $1 - \frac{900}{9600} > 0.9$. Moreover, ω_{DP}^* should be less than 1.0, corresponding to *DP*, which goes to zero. Thus, putting $2 - \omega_{DP}^* \cong 1$ in Equation (A8), we have

$$\frac{\Delta CB_{nor}(t,T_r)}{\Delta t} \cong \frac{k_{DP}(T_r)}{4}$$
(A9)

Inserting Equation (A9) into the definition of the Arrhenius shift factor a_{TDP} , given by Equation (A5), we can relate the *DP* activation energy E_{DPa} to this slope:

$$E_{DPa} = \frac{RT_oT_r}{T_o - T_r} \times ln \left[\frac{4 \times \frac{\Delta CB_{nor}(t, T_r)}{\Delta t}}{k_{DP}(T_o)} \right]$$
(A10)

Appendix B: Noise-to-Signal Ratio Statistical Analysis of the X-Ray Dating

In this section, we analyze the robustness of the X-ray prediction of the aging times calculated from Equation (8). This deeper study can be done with the theory of regression because the best relationship between the X-ray aging time and the ¹⁴C aging time should be linear before saturation occurs, and a novel analysis of its results considers a suitable "noise" and a noise-to-signal ratio and its geometrical representation. We first consider the prediction of the ¹⁴C data from the X-ray and then the opposite. We exclude by the following analysis sample FII, characterized by a HR dating, because it has been used to calculate linen aging quantities used in the X-ray dating.

Let *x* be the X-ray aging time reported in the third column of Table 3, and let *y* be the central/median value of ${}^{14}C$ aging time reported in the second column of Table 3.

Figure A1 shows the three scatter plots and their best-fit regression lines between (i) the central values of *x* vs. *y*; (ii) the +2 standard deviations bound (2σ) of *x* vs. *y*; and (iii) the -2 standard deviation (-2σ) bound of *x* vs. *y*. We notice that a regression line *y* = *ax* fits the data very well. We propose to estimate the robustness of the linear regression lines between *y* and *x* obtainable from

Table 3 by using a suitable noise-to-signal ratio (NSR) and its geometrical representation, the latter borrowed from the optimum design of deep-space 2-hop radio links [27].

Two variables, *y* and *x*, linked by a regression line y = ax, where *a* is the slope, are perfectly correlated if r = 1 and are not biased if a = 1, in other words, if the regression line is y = x (45° line, a = 1) and all *y* values lie along the line (r = 1). For the central value of Figure A1 (black line), a = 0.9996 and r = 0.9959. If one or both of these conditions are not met, we can take the differences between the actual regression line—with $a \neq 1$ and r < 1—and the ideal line as "noise". Defining the variances of *x* and *y* as s_x^2 and s_y^2 , the variance (power) of the noise due to $a \neq 1$ (regression noise) is given by the following [28]:

$$N_a = (a - 1)^2 s_x^2$$
 (A11)

The power of the noise due to all the *y* values not lying along the line is given by the following:

$$N_r = (1 - r^2) s_y^2$$
(A12)

Therefore, the total noise-to-signal ratio is given by the sum of the two unconnected single ratios:

$$NSR = \frac{N_a}{s_x^2} + \frac{N_r}{s_x^2} = NSR_a + NSR_r = (a-1)^2 + (1-r^2)\frac{s_y^2}{s_x^2}$$
(A13)

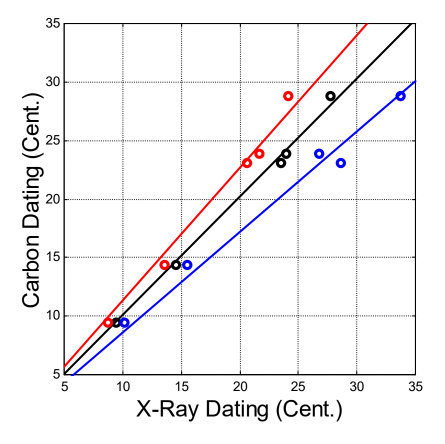


Figure A1. Scatter plots and their best-fit regression lines between (i) the central values of *x* vs. *y* (black); (ii) the $+2\sigma$ values of *x* vs. *y* (blue); and (iii) the -2σ of *x* vs. *y* (red).

Because $r^2 s_y^2 = a^2 s_x^2$, Equation (A13) can be written as follows:

NSR =
$$(a-1)^2 + \frac{(1-r^2)}{r^2}a^2$$
 (A14)

Equation (A13) graphically describes a circle with radius $\sqrt{\text{NSR}} = \sqrt{\text{NSR}_a + \text{NSR}_r}$ [27]. Therefore, in the first Cartesian quadrant, $\sqrt{\text{NSR}_a}$ and $\sqrt{\text{NSR}_r}$ for a constant NSR = NSR_o lie in a quarter of a circle, as shown in Figure A2 for the three cases of the variable *x*. All values inside the circle with radius $\sqrt{\text{NSR}_o}$ have NSR < NSR_o. The origin has NSR = 0; therefore, *a* = 1 and *r* = 1. The reciprocal of NSR is the signal-to-noise ratio, SNR, which becomes infinite at the origin. As discussed in Reference [27], among other features not of interest here, the NSR allows a graphical representation that clearly shows how the partial NSR_r and NSR_a contribute to the total NSR.

Figure A2 shows the results obtained using the central values of the X-ray aging time and the 95% bounds (-2σ , 2σ) reported in Table 3: (i) only for the central values the NSR is dominated by NSR_r because $\sqrt{\text{NSR}_a} < \sqrt{\text{NSR}_r}$; (ii) the central values of Table 3 give the lowest (highest) NSR (SNR), while the $\pm 2\sigma$ values gives similar and markedly larger NSRs. For the central value, the slope is in practice 1 (i.e., a 45° line); therefore, NSR_a \approx 0, as Figure A2 shows.

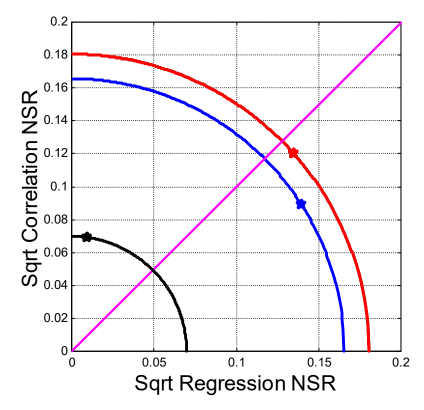


Figure A2. From X-ray to ¹⁴C measurements: Square root of the noise-to-signal ratio (NSR) as a function of the square root of the regression NSR_{*a*} (abscissa) taken as the absolute value |a - 1| and the square root of the correlation NSR_{*r*} (ordinate): The black line is the central values of column 3 in Table 3. The blue line is $+2\sigma$. The red line is -2σ . The 45° line shows where NSR_{*r*} = NSR_{*a*}.

The point which graphically represents the central value NSR on the black circle in the first Cartesian quadrant is very close to the vertical axis. Conversely, the $\pm 2\sigma$ NSRs are dominated by NSR_a. This finding is related to the saturation with the aging time of the natural degradation of cellulose towards the maximum allowed values, which causes an increase of the uncertainty interval as a function of *CB*_{nor}. The different widths of $\pm 2\sigma$ and -2σ uncertainty intervals for large *CB*_{nor} values close to 0.98 have the same origin. The association of an asymmetric $-2\sigma_r + w\sigma$ uncertainty interval at the central value, with w < 2, in case of very large aging times (*CB*_{nor} close to 0.98) would partially

reduce the effect of CB_{nor} saturation on the statistical noise which, in these cases, is dominated by NSR_r. In any case, the central values of the X-ray dating minimizes the NSR and is practically not polarized, with a negligible NSR_a because $a \cong 1$. Therefore, it gives an accurate estimate of the aging time, at least in terms of comparison with the ¹⁴C dating.

Now, let us consider the results the prediction of the X-ray aging time from the ¹⁴C data of Table 3. Figure A3 shows the regression lines fitted to the data, while Figure A4 reports the NSR Cartesian quadrant. The ¹⁴C data are less scattered around the central values, as Table 3 reports and Figure A3 clearly shows in the estimate of the X-ray prediction bounds.

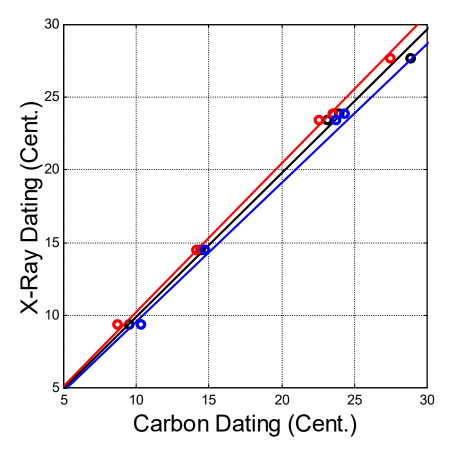


Figure A3. Scatter plots and their best-fit regression lines between: (i) the central values (black); (ii) the $+2\sigma$ values (blue); and (iii) the -2σ values(red).

We can now see that the central values give the same NSR of Figure A2. This means that the prediction works with the same precision in both directions. Moreover, contrarily to Figure A2, the three NSRs in Figure A4 are dominated by NSR_a; therefore, there is not a large polarization of the statistics as is the case with the $\pm 2\sigma$ and $\pm 2\sigma$ bounds reported in Figure A3. Therefore, if we consider the central values of Table 3, we can state that the two methods of estimating the aging time, namely the ¹⁴C and the X-ray, are practically indistinguishable. If we consider $\pm 2\sigma$ bounds, the X-ray dating can be affected by statistical polarization for large *CB_{nor}* values close to 1 due to the saturation of the cellulose degradation for large aging times. This finding leads to reliable X-ray dating only when the linen fabrics have suffered a natural aging at an average room temperature of 21–23 °C for not more than 25–30 centuries. For smaller average room temperatures, linen fabrics also older than 25–30 centuries could be reliably dated by the proposed X-ray method.

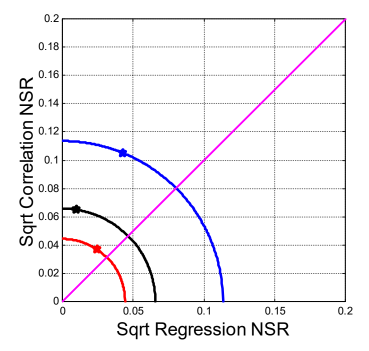


Figure A4. From ¹⁴C to X-ray measurements: Square root of the noise-to-signal ratio (NSR) as a function of the square root of the regression NSR_{*a*} (abscissa) taken as the absolute value |a - 1| and the square root of the correlation NSR_{*r*} (ordinate): The black line is the central values of column 2 in Table 3. The blue line is $+2\sigma$. The red line is -2σ . The 45° line shows where NSR_{*r*} = NSR_{*a*}.

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