CHAPTER ONE
Chapter one

1.1 Introduction

Measuring of entrance and exit dose is most important in radiation therapy because of the effect of radiation in the skin of the patient, and Prediction of skin reactions requires knowledge of the dose at various depths in the human skin. Using thermoluminance dosimeters (TLD) of different thicknesses, the dose can be extrapolated to the surface and interpolated between the different depths.

The thermo-luminescent (T.L.) phosphoric material such as LiF(Mg, Cu and P) which are usually named (TLD100H) has shown great interest of scientists in the field of radiation dosimetry due to its very favorable dosimetric characteristics (Fung 2004). This material was first introduced in the powder form by Nakajima et al. in 1978 then the development continued and commercialized in 1990 as LiF:Mg.Ti (TLD100) and the LiF(Mg,Cu,P) T.L. phosphors which are tissue-equivalent. The TLD100H is considered as outperforms due to its very higher sensitivity, by a factor of greater than 25, and a dose detection threshold of less than 1 mGy (Moscovitch 1999). Therefore TLD has been used in the field of radiation dosimeter and correlation of dose to biological effects (Ayad 2000).

One of the most promising realms being under focused by scientists in the nanocomposites materials which have many applications in the field of medical engineering, electronics, electrochemical, biosensors, radiation detection/measurements and optical/display systems that is for their wide range of absorption frequencies and ultra fast response behaviors (1). Nanocomposites martials categorized under nanotechnology which can be defined as the science dealing with the construction and use of functional structures designed form atomic or molecular
scale which at least one characteristic dimension measurement in nanometers (Kensuke et al., 2000; Carl, 2002).

All metallic nanoparticles (Gold, Silver, Platinum…etc) exhibit unusual electrical, magnetic, optical and electrochemical properties due the near-free conduction electrons, which are dependent on their size, surface Plasmon, surface free energy and surface area, as well as on the surrounding dielectrics (Stakeev and Kustov, 1999).

The recent studies on polymer/silver nanocomposites revealed that the properties of silver within it is composites answer many potential applications in optical waveguides, optical switches, molecular identification, oxidative catalysis, antimicrobial effects, etc.

These mentioned properties are dependent on the particle size, shapes and the method of composites synthesis. Varieties of synthesis method have been developed, including reduction from metallic salts, ultrasonic irradiation technique, ion implantation, and thermal process and microwave technique. The synthesis of silver nanocomposites by using γ- irradiation has been mentioned by Ali et al., (2007). The radiation measurements in this study will be depend on the optical density measurement and optical absorbance measurement using optical densitometer and UV-visible spectroscopy accordingly. Also the radiation effect on the Silver/ PVA composites will be shown out of SEM studies. The expected x-ray or γ exposure on the silver/PVA composites film will be as color change to yellow or golden due to presence of silver while the expected change on PVA will be as degradation of the PVA chain.
1.2 **Problem of the study**

The following study will help to establish a local in vivo dosimeter system to measure the entrance and exit dose with local available materials in Sudan radiotherapy centers. Also the private clinics lacking the frequent radiation exposure assessment tools, hence it will be as a fast, easy and cheap assessment tool with a presence of densitometer which is the cheapest equipment used in the radiation exposure assessment.

1.3 **Significant of the study**

Ionizing radiation has significant capabilities to induce chemical and physical change in the exposed materials. Studied on irradiated polymers show many advantage of induced radiation changes on which can be quantify and qualify to deduce the amount of radiation dose and exposure. This method relative to other methods such as film badge is so simple and easy. The advantage includes no developer, no fixer, no other effecting factors, the process occurred at a solid-site condition, fast method and inexpensive, environmental friendly and controllable acquisition (6).

1.4 **Scope of the study**

Ionizing radiation has significant capabilities to induce chemical changes in the exposed materials.

So the following study will highlights and shows the exposure dose based on the PVA/Ag composites change due to irradiation by using densitometer and UV-visible spectroscopy.
1.5 Hypothesis of the study

The researcher expect that Sudanese radiation therapy centers have high exposure dose to the skin of the patients during treatment with radiation which is incompatible with the policy of radiation therapy system. Also there is lack in the facilities for assessment and measuring the in vivo dosimetry especially entrance and exit dose.

1.6 Objectives of the study

- To measure entrance and exit dose to the patients during radiotherapy treatment.
- To evaluate the PVA/Ag composites film as measuring tool in radiotherapy especially in in-vivo dosimetry depending on optical density or absorption coefficient.
- To identify the radiation effect on the PVA/Ag composites film.
- To compare the PVA/Ag composites film exposure data with TLD.
- To determine the absorption coefficient of the phantom depending on the entrance and the exit dose.

1.7 Thesis outline

The backbone of the following thesis will be formed of five chapter. Chapter 1 will deal with the general introduction about the research background, scope of the study, problem statement, Hypothesis of the study and the objectives of the study. Chapter 2 will deal with literatures review cover the theoretical background and previous studies. Chapter 3 will deal with the methodology of the study, including materials, samples preparation, equipment and measurement. Chapter 4 will cover the results. And chapter 5 will cover discussion, conclusion, recommendations, references and the researcher bio data.
CHAPTER TWO
Chapter two

Literature Review

2.1 Theoretical background

2.1.1 Ionizing radiation

Ionizing radiation is a broad energetic spectrum of electromagnetic waves or high velocity atomic or subatomic particles. The radiation can be categorized according to their ability to ionize the media. Non-ionizing radiation is electromagnetic radiation that does not have sufficient energy to remove an electron of the atom. The various types of non-ionizing radiation are ultra violet (UV), visible light, infrared (IR), microwaves (radio and television), and extremely low frequency (ELF, or as they called EMF or ELF-EMF). Ionizing radiation is electromagnetic radiations, such as X-rays, γ-rays and charged particles (electrons, α-particles and β-particles) which possess sufficient energy to ionize an atom by removing at least an orbital electron. According to the 1996 European Guideline of the European Atomic Energy Community (EURATOM), electromagnetic radiation with a wavelength of 100 nm or less is considered as ionizing radiation which is corresponds to ionizing potential of 12.4 eV or more (Smith, 2000). The ionization potential is dependent on the electronic structure of the target materials and generally in the order of 4 – 25 eV.

The International Commission of Radiation Units (ICRU) has subdivided the ionizing radiation into direct and indirect ionizing radiation, based on the mechanisms by which they ionized the atom. Direct ionizing radiations are fast charged particles, such as alpha particles, electrons, beta particles, protons, heavy ions, and charged mesons, which transfer their energy to the orbital electron directly and ionize the atom by means of Columbic force interactions along their track. Indirect ionizing radiations are uncharged quantum, such as electromagnetic radiations (X-rays and γ-
rays), neutrons, and uncharged mesons, which undergo interactions with matter by indirectly releasing the secondary charged particles which then take turn to transfer energy directly to orbital electrons and ionize the atom. Some properties of ionizing radiation are shown in Table 2.1

**Table 2.1 shows the properties of different ionizing radiation**

<table>
<thead>
<tr>
<th>Characteristics</th>
<th>Alpha</th>
<th>Proton</th>
<th>Beta or electron</th>
<th>Photon</th>
<th>Neutron</th>
</tr>
</thead>
<tbody>
<tr>
<td>Symbol</td>
<td>^2_2α or He$^{+2}$</td>
<td>^1_1 p or H$^+$</td>
<td>^-1_e or β</td>
<td>γ- or X-rays</td>
<td>^0_0 n</td>
</tr>
<tr>
<td>Charge</td>
<td>+2</td>
<td>+1</td>
<td>-1</td>
<td>Neutral</td>
<td>Neutral</td>
</tr>
<tr>
<td>Ionization</td>
<td>Direct</td>
<td>Direct</td>
<td>Direct</td>
<td>Indirect</td>
<td>Indirect</td>
</tr>
<tr>
<td>Mass (amu)</td>
<td>4.00277</td>
<td>1.007276</td>
<td>0.000548</td>
<td>-</td>
<td>1.008665</td>
</tr>
<tr>
<td>Velocity (m/s)</td>
<td>6.944 x10^6</td>
<td>1.38 x10^7</td>
<td>2.82 x10^8</td>
<td>2.998 x10^8</td>
<td>1.38 x10^7</td>
</tr>
<tr>
<td>Speed of light</td>
<td>2.3%</td>
<td>4.6%</td>
<td>94%</td>
<td>100%</td>
<td>4.6%</td>
</tr>
<tr>
<td>Range in air</td>
<td>0.56 cm</td>
<td>1.81 cm</td>
<td>319 cm</td>
<td>820 m</td>
<td>39.25 cm</td>
</tr>
</tbody>
</table>

1 atomic mass unit (amu) = 1.6 x 10^{-27} kg.
Speed of light $c = 3.0$ x 10^8 m/sec.

**2.1.2 Radiation sources**

The sources of ionizing radiation can be divided into two categories namely as natural and man-made sources. The first natural sources are cosmic radiation which is the radiation coming from outside our solar system as positively charged ions (protons, alpha, heavy nuclei) and interact with atmosphere to produce secondary radiations such as x-rays, muons, protons, alpha particles, pions, electrons and neutrons. The second natural sources are external terrestrial sources which represent the radioactive materials found naturally in the earth crust, rocks, water, air and vegetation. The
major radionuclides found in the earth crust are Potassium-40, Uranium-235, and Thorium-210. The main sources of man-made radiations that expose the public are in the form of medical diagnostic X-ray, radiation therapy, nuclear medicine and sterilization. The common radioactive sources are $^{131}$I, $^{99m}$Tc, $^{60}$Co, $^{192}$Ir, $^{90}$St and $^{137}$Cs. Other man-made sources exemplar in occupational and consumption products that originated in mines, combustible fuel (gas, coal), ophthalmic glasses, televisions, luminous, watch’s dial (tritium), X-rays at the airport (scanner), smoke detectors (Americium-241) and fluorescent lamp starters, nuclear fuels, nuclear accidents and nuclear weapons. The yield of artificial sources is represented either as quantum in X-rays and $\gamma$-rays or as high energy particles such as beta particles ($\beta$), alpha particles ($\alpha$), neutrons and electrons (Smith, 2000). Most of the types of radiation source discussed above are used in medicine, industry, and research. Today the most common radiation sources used is $^{60}$Co, which is an artificial source of $\gamma$ radiation and linear accelerators for photon and electron beams having potential energy ranging from 0.3-10 MeV to 20 MeV. Table 2.2 shows the radiation sources commonly used in industry and research.

Table 2.2 shows the common sources of ionizing radiation. (Smith, 2000)

<table>
<thead>
<tr>
<th>Category</th>
<th>Source</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nuclear power</td>
<td>$^{235}$U fission products, $^{90}$Sr, $^{137}$Cs</td>
</tr>
<tr>
<td>Occupational exposure</td>
<td>X-ray, isotopes for $\gamma$-rays</td>
</tr>
<tr>
<td>Weapons tests</td>
<td>$^{233}$U, $^{239}$Pu, fission products</td>
</tr>
<tr>
<td>Every day sources</td>
<td>Coal, Tobacco and Air-travel</td>
</tr>
<tr>
<td>Medical tests &amp; treatment</td>
<td>X-ray, $\gamma$-rays &amp; electrons</td>
</tr>
<tr>
<td>Cosmic rays</td>
<td>Protons, electrons, neutrons</td>
</tr>
<tr>
<td>Food</td>
<td>$^{40}$K, $^{137}$Cs, $^{14}$C and $^{131}$I</td>
</tr>
<tr>
<td>Rocks &amp; building</td>
<td>$^{235}$U, $^{238}$U, and $^{232}$Th</td>
</tr>
<tr>
<td>Atmosphere</td>
<td>$^{222}$Rn and $^{137}$Cs</td>
</tr>
</tbody>
</table>
2.1.3 γ- radiation sources

γ-rays are produced by the nuclear transitions that occur within the nuclei of radioactive elements. The emitted photons are mono-energetic with specific energy to the isotope from which they originate. By far the most commonly employed radioactive isotope for γ-rays is cobalt-60 (60Co), an isotope with a half-life of 5.272 years. 60Co emits two γ-photon s of equal intensity at 1.17 and 1.33 MeV. It is produced in nuclear reactors by a neutron-capture reaction involving 59Co. Due to the long half-life, high penetrating power and ease of production, 60Co sources have become sources of choice in both industrial and research institutions. The activity of the sources prepared in nuclear reactors can be made high up to 40 Ci/g (1.5 TBq/g), however sources with activity from 1 to 5 Ci/g are typically common for use. Over 80% of the 60Co produced world-wide is manufactured by the Canadian company, Ontario Hydro and marketed by another Canadian company, MDS Nordion.

60Co radioisotope decays to stable nickel-60 by a nuclear transition (β-decay) in which a neutron is converted into a proton via the emission of a β-particle with energy of 312 keV and two γ-photon s, one of 1.17 MeV and another of 1.33 MeV. The decay scheme of this isotope is illustrated in Figure 2.1 (Choppin et al., 1995; Attix, 1986).
Another frequently used $\gamma$-ray source is cesium-137, a fission product from nuclear reactors. The energy of the emitted photon is 662 keV and the half-life is 30.17 years. Nuclear reactors themselves are potential sources of $\gamma$-rays.

### 2.1.4 $\gamma$-radiation interaction with matter

When photons of $\gamma$-radiation interact with matter, they undergo attenuation and hence lose their energy and intensity by the process of photoelectric absorption, Compton inelastic scattering, pair production, and Rayleigh elastic scattering (Evan, 1952).

### 2.1.5 Photoelectric absorption

Absorption of $\gamma$-rays occurs when the $\gamma$-ray photon is absorbed by an electron resulting in ejection of the electron from the inner shell of the atom and ionization of atom take place. Subsequently, the ionized atom returns to the neutral state with the emission of characteristic X-ray of the atom shown in Figure 2.2. This subsequent emission of lower energy photons is generally absorbed and does not contribute to (or hinder) to the secondary ionization. Photoelectron absorption is the dominant process for $\gamma$-ray absorption up to energies of about 500 keV. The photoelectric absorption process predominates for photons in the low energy range between 10 keV and 200...
keV. This is the phenomenon explained by Einstein in 1905, in which an incident photon gives up all its energy $h\nu$ to a bound electron, usually K shell (90%), where subsequently part of the energy is used to overcome the electron binding energy $E_B$ and the extra energy is converted as kinetic energy $K_E$ of the photoelectron. This can be expressed in equation (1.1).

$$h\nu = K_E + E_B$$

(2.1)

The atom that is left in an excited state will emit fluorescent x-rays or Auger electrons. The characteristic X-rays may escape especially for high-energy photons and high atomic number of absorbing material unless the absorber is thick enough to stop the $\gamma$-rays. The ranges of the Auger electrons are short and locally absorbed. The cross-section for photoelectric effect in K shell of an atom with atomic number $Z$ for photon energy $h\nu$ is given by equation (2.2), from which we could deduce that the photoelectron absorption is dominant for atoms of high atomic numbers and for photon of low energy.

$$\sigma_K = \varphi_0 4\sqrt{2} \left( \frac{m_0c^2}{h\nu} \right)^{3/2} \frac{Z^5}{137^4}$$

(2.2)

where $\varphi_0 = \pi r_0^2$ and $r_0 = e^2 / m_0 c^2 = 2.818 \times 10^{-13} m$ is the classical electron radius, $m_0$ is the rest mass of electron and $c$ is the speed of light. $m_0 c^2$ is the rest energy of the recoil electron, according to mass-energy equation proposed by Einstein in 1905 in the relativity theory.
2.1.6 Compton Scattering

Compton scattering process also known as an incoherent scattering, occurs when the incident photon ejects free or weakly bonded electron from an atom and a photon of lower energy is scattered from the atom. Relativistic energy and momentum are conserved in this process and the scattered $\gamma$-ray photon has less energy and therefore greater wavelength than the incident photon shown in Figure 2.3. Compton scattering is important for low atomic number specimens. At energies of 100 keV - 10 MeV the absorption of radiation is mainly due to the Compton effects (McGervey, 1983). The change in wavelength of the scattered photon is given by equation (2.3)

$$\frac{c}{v} - \frac{c}{v_o} = \lambda - \lambda_o = \frac{h}{mc(1-\cos \theta)}$$  \hspace{1cm} (2.3)
where \( \lambda \) is the wavelength of the incident photon, \( \lambda_0 \) the wavelength of the scattered photon, \( m \) is the mass of the electron and \( \theta \) the angle of scattering for the photon. On rearranging, the above equation we get the following equation (1.4)

\[
hv = \frac{h}{1 + \alpha(1 - \cos \theta)}
\]  

(2.4)

Where \( \alpha \) is the ratio of the energy of the photon to the rest energy of the electron i.e. \( \alpha = h\nu / m_0c^2 \). The kinetic energy \( T \) for the recoil electron is given by equation (1.5)

\[
T = h\nu \frac{\alpha(1 - \cos \theta)}{1 + \alpha(1 - \cos \theta)}
\]  

(2.5)

And the scattering angle of the electron is given by equation (1.6)

\[
\cot \phi = (1 + \alpha) \tan \frac{1}{2} \theta
\]  

(2.6)

Figure 2.3  Schematic diagram of Compton scattering for \( \gamma \)-radiation resulting in ionization and scattering of the incident photon with less energy
2.1.7 Pair Production

The production of a positive and negative electron pair (pair production) is a process that can take place in the vicinity of the nucleus field of an atom or an electron field. Absorption of photons through the mechanism of pair production can occur when the energy of an incident photon is greater than twice the rest mass of an electron, i.e. $2m_0c^2 = 1.022$ MeV (Johns and Cunningham, 1983). During pair production interactions, a photon has its energy converted to an electron–positron pair. The positron so produced interacts with matter by ionizing and exciting atoms through the same processes as electrons, thus losing energy and being brought to rest. At this point, the positron combines with an electron in an annihilation process producing two photons with energy equal to 0.511 MeV as shown in Figure 2.4.

Pair production is an absorption process in which a photon in the field of nucleus produces an electron-positron pair, where the total kinetic energy is equal to the energy of photon minus the rest energy of the two particles, which have been created as shown in equation (2.7), thus the photon energy must be greater than 1.02 MeV for the interaction to take place. The electron and positron do not necessarily receive equal energy, but their average energy is given by equation (2.8).

$$h\nu = (m_0c^2 + T_e) + (m_0c^2 + T_+$$)

$$2 \nu = \frac{h\nu - 1.022}{2} \text{ MeV}$$

The cross-section of the pair production in the field of nucleus is given by equation (2.9)

$$\sigma_{pp} = \frac{1}{137} r_0^2 Z^2 \left[ \frac{28}{9} \ln \left( \frac{2h\nu}{m_0c^2} \right) - \frac{218}{27} \right]$$

The triplet production process is similar but the interaction takes place with one of the atomic electrons which receives sufficient energy to be set free. It occurs when the
incident photon have an energy of $4m_0c^2$, i.e. it implies both the pair production at the nucleus level plus triplet production. The total kinetic energy is equal to the energy of photon minus the rest energy of the three ejected particles as given by equation (2.10), from which the average kinetic energy can be deduced by equation (2.11) (Motzet al., 1969)

$$hv = (m_0c^2 + T_e) + (m_0c^2 + T_M) + (m_0c^2 + T_P)$$

$$\bar{T} = \frac{hv - m_0c^2}{3} \text{ MeV}$$

**Figure 2.4** Schematic diagram of Pair Production process for $\gamma$-radiation being interfered in the nucleus field and orbital electron to produce triplet particles.

### 2.1.8 Raleigh scattering

The coherent Rayleigh scattering by an atom is predominant for photons at low energy range from 1 keV to 100 keV. The Rayleigh scattering is a process in which a photon is deflected by a bounded electron of the atom and photon going off in
different directions with no loss in energy. The atomic system may recoil as a whole under impact without the atom being ionized or excited. The probability of this process is large only for low energy photons and high atomic number material. The differential cross-section of the coherent scattering of photon at deflection angle $\theta$ by a bounded electron is given by equation (2.12)

$$d\sigma_{coh} = \frac{1}{2} f_0^2 (1 + \cos^2 \theta) \left[ \frac{F(\alpha, \theta, Z)^2}{Z} \right] d\Omega$$

(2.12)

where $F(\alpha, \theta, Z)$ is the atomic form function which varies from zero at large angle to $Z$ at the smaller angle.

2.1.9 $\gamma$-ray attenuation coefficients

In general the characteristic of radiation interaction with matter is that each individual photon is absorbed or scattered from the incident beam in a single event. The photon number removed $\Delta I$ is proportional to the thickness traveled through $\Delta x$ and the initial photon number $I_0$, i.e. $\Delta I = -\mu I_0 \Delta x$, where, $\mu$ is a constant proportionality called the attenuation coefficient. In this case, upon integrating, we have the following equation (2.13)

$$I = I_0 e^{-\mu x}$$

(2.13)

The attenuation coefficient is related to the probability of interaction per atom, i.e. the atomic cross section $\sigma_a$ is given by equation (2.14)

$$\mu = \frac{\sigma_a N_A \rho}{A}$$

(2.14)
Where \( A \) is the mass number and \( N_A \) the Avogadro’s number \((6.022 \times 10^{23}\text{mol}/1)\), Table 2.3 briefly summarized the entire \( \gamma \)-radiation photon interactions with their possible energies required to initiate the reactions (Smith, 2000; Siegbahn, 1965).

<table>
<thead>
<tr>
<th>Process</th>
<th>Type of interaction</th>
<th>Other names</th>
<th>Approximate ( E ) of Maximum importance.</th>
<th>( Z ) dependence</th>
</tr>
</thead>
<tbody>
<tr>
<td>Photoelectric</td>
<td>With bonded electrons, all ( E ) given to electron</td>
<td>Rayleigh electron, resonance scattering, Thomson scattering</td>
<td>Dominant at low ( E ) ((1\text{-}500)\text{KeV}, \text{cross section decrease as} \ E \text{ increase})</td>
<td>( Z^3 )</td>
</tr>
<tr>
<td>Scattering from electrons coherent</td>
<td>With bond atomic electron, with free electrons</td>
<td>Compton scattering</td>
<td>(&lt;1\text{MeV and greatest at small angles. Independent of energy})</td>
<td>( Z^2, Z^3 )</td>
</tr>
<tr>
<td>Incoherent</td>
<td>With bond atomic electron, with free electrons</td>
<td>Elastic Pair production</td>
<td>(&lt;1\text{MeV least at small angle. Dominate in region of 1 MeV, decreases as} \ E \text{ increase})</td>
<td>( Z )</td>
</tr>
<tr>
<td>Pair Production</td>
<td>In Coulomb field of Nucleus</td>
<td>Triplet production inelastic pair production. Nuclear potential scattering</td>
<td>Threshold ( \sim1\text{MeV, } E&gt;5\text{MeV}. \text{Increase as} \ E \text{ increase.})</td>
<td>( Z^2 )</td>
</tr>
<tr>
<td>Pair production Delbruk scattering</td>
<td>In coulomb field of electron &amp; nucleus</td>
<td></td>
<td>Threshold at 2 \text{MeV} increases as ( E ) increases. Real Max ( &gt;) imaginary, below 3 ( \text{MeV(both increase as} \ E \text{ increases)})</td>
<td>( Z )</td>
</tr>
</tbody>
</table>

**2.1.10 \( \gamma \)-radiation interaction with molecules**

The essence of \( \gamma \)-radiation interaction with molecules and change the physical and chemical characteristics of the formed compound is ascribed to the amount of energy being transferred, which will create ion, free radicals and excited molecule. Such interaction process is termed ionization and excitation of the molecules, which can cause chemical changes to the irradiated molecule. This is due to the fact that all
binding energy for organic compound in the range of 10 – 15 eV. In case of low transferred energy by photon, the molecule undergoes excitation state before returning to the rest state by emitting X-ray photons or break down to release free radicals which in turn undergoes polymerization.

The ejected electron from the irradiated molecule (A\(^+\)) is subjected to the strong electric field of the formed positive charge. Therefore the recombination is a frequently occur, either during irradiation or after the end of irradiation to create energetic molecule (A\(^{**}\)). Such highly energetic excited molecule will break down into free radicals and new molecule (Denaro, 1972). The fundamental of this reaction can be shown in the following scheme Figure 2.5.

![Reaction Scheme](image)

Figure 2.5 shows the expected irradiation results of the organic molecules, where R, and S, are free radicals and M and N are molecular products.
2.1.11 Polymer

Polymer is a Greek word contain of two word poly +mer and they mains many parts, polymer is a large molecule, or macromolecule, composed of many repeated subunits. Because of their broad range of properties (Painter et al 1997), both synthetic and natural polymers play an essential and ubiquitous role in everyday life. (McCrum et al 1997), Polymers range from familiar synthetic plastics such as polystyrene to natural biopolymers such as DNA and proteins that are fundamental to biological structure and function. Polymers, both natural and synthetic, are created via polymerization of many small molecules, known as monomers. Their consequently large molecular mass relative to small molecule compounds produces unique physical properties, including toughness, viscoelasticity, and a tendency to form glasses and semi crystalline structures rather than crystals.

2.1.12 Polymer properties

Polymer properties are broadly divided into several classes based on the scale at which the property is defined as well as upon its physical basis. (S.A. Baeurle, 2009). The most basic property of a polymer is the identity of its constituent monomers. A second set of properties, known as microstructure, essentially describe the arrangement of these monomers within the polymer at the scale of a single chain. These basic structural properties play a major role in determining bulk physical properties of the polymer, which describe how the polymer behaves as a continuous macroscopic material. Chemical properties, at the nano-scale, describe how the chains interact through various physical forces. At the macro-scale, they describe how the bulk polymer interacts with other chemicals and solvents.

2.1.13 Polymerization

Radiation polymerization is a process in which the free radicals interact with the unsaturated molecules of a low molecular unit known as monomer to form high
molecular mass polymer or even with different monomers to produce crosslink polymer. The formed polymer can be in different forms called homopolymer and copolymer depending on the monomer compositions link together.

Radiation-induced polymerization process can be achieved in different media whether it is liquid or solid unlike the chemical polymerization which can only accomplished in aqueous media. It is also temperature independent. Radiation polymerization often continues even after removing away from the radiation source. Such condition is known as post-polymerization (Lokhovitsky and Polikarpov, 1980). Since radiation initiation is temperature independent, polymer can be polymerized in the frozen state around aqueous crystals. The mechanism of the radiation induced polymerization is concerning the kinetics of diffusion-controlled reactions and consists of several stages: addition of hydroxyl radicals and hydrogen atoms to carbon-carbon double bond of monomer with subsequent formation of monomer radicals; addition of hydrated electrons to carbonyl groups and formation of radical anion of a very high rate constant and the decay of radicals with parallel addition of monomer to the growing chain.

### 2.1.14 Cross linking

The process of crosslink occurs due to interaction between two free radical monomers which combine to form intermolecular bond leading to three dimensional net of cross linked highly molecular polymer, more likely dominate in unsaturated compound or monomer. The cross linked polymer show strong mechanical strength and high thermal resistance.
2.1.15 Radiation grafting

Radiation grafting is a process in which active radical sites are formed on or near the surface of an exciting polymer, followed by polymerization of monomer on these sites. Grafting is accompanied by homopolymerization of the monomer; the material to which the monomer is grafted is described as the backbone, trunk or support. Radiation grafting is used to modify the polymers texture such as film, fibers, fabrics and molding powders.

The process of grafting can be expressed as follow; suppose the polymer A is exposed to γ-rays, thus the active free radical sites A* created randomly along the polymer backbone chain, this free radical initiate a free radical on the monomer B then undergoes grafting polymerization at that active sites. The extension of the attached monomer B upon the base polymer A is termed as the degree of grafting (DOG) which refers to the mass of the grafted polymer as a percentage of the mass of the original base polymer. Such process can be expressed in schematic Figure 2.6.

Figure 2.6 shows the schemes for grafting process for polymer A with monomer B using γ-radiation.
2.1.16 Absorption of light and UV-Visible spectrophotometry

2.1.16.1 Optical absorption

The transmittance of light across the absorber accompanied with absorption of the light waves. Such absorption depends on the wavelength of the light, thickness of the absorber or the transmitted media and the nature of the media. Beer-Lambert Law, successfully determined the absorption coefficient of the media as follows; supposing a light beam \( I \) is being incident to an object whose area is \( a \), \( \alpha \) is its absorption coefficient, then the fraction absorption \( dl \) of light by this absorber in a thickness \( dx \) is given by the following equation (2.15)

\[
dl = -\alpha I dx
\]  

(2.15)

or

\[
\frac{dl}{I} = -\alpha dx
\]  

(2.16)

The integrating of this equation (2.16) gives the absorbance \( A \) as in equation (2.17). In this case the light intensity \( I \) changed from 0 to \( I \) by crossing a distance from \( x = 0 \) to \( x = x \) in the absorber.

\[
\int_{l=0}^{l=I} \frac{dl}{I} = -\alpha \int_{x=0}^{x=x} dx
\]

\[
\ln I / I_0 = -\alpha x / I_0
\]

\[
\ln(I) - \ln(I_0) = -\alpha x
\]

\[
\ln \frac{I}{I_0} = -\alpha x
\]

\[
\log \frac{I}{I_0} = e^{-\alpha x} = A
\]  

(2.17)
From equation (2.17) of absorbance $A$, the absorption coefficient $\alpha$ can be deduced in equation (2.18).

$$\log \frac{I}{I_0} = \alpha x$$

$$\frac{e \times A}{x} = \alpha$$

(2.18)

As the value of $\log \frac{I}{I_0} = A$ and $e$ is 2.303, then the absorption coefficient can be given by equation (2.19)

$$\alpha = \frac{2.303 \times A}{x}$$

(2.19)

Where $x$ is the distance crossed by the light or can be the sample thickness in case of UV-visible spectroscopy. The corrected absorption value known as the molar absorptivity ($\varepsilon$) is used for the comparison between different spectra of different compounds and can be expressed by the following equation:

$$\varepsilon = \frac{A}{cl}$$

(2.20)

Where $A = $ absorbance, $c = $ sample concentration in moles/liter and $l = $ path length through the cuvette in cm.

2.1.16.2 Mechanism of absorption process

The absorption processes of microwaves in polymer, generally accompanied with energy reduction in the transition wave. Such reduction is due to the optical absorption constant of the media $\alpha$ which come in the range of $10^5 - 10^6$ cm$^{-1}$ (Richard, 1988). The variation of the absorption on the transition involves a photon only (direct transition), or that involves both a photon and a phonon (indirect
transition). As the photon energy drops below the band gap energy, the absorption constant decreases by many orders of magnitude.

The energy levels are created if imperfections are present in the polymers. They lie in forbidden gap. At energy less than the band gap energy it is still possible to excite electrons to the conduction band from imperfection levels occupied by electrons. Electrons also can be excited from the valence band to unoccupied imperfection levels. Each process will give rise to optical absorption. When the photon energy is less than the energy required to make a transition from the imperfection level to one of the bands, this absorption in turn comes to an end. The corresponding absorption constant $\alpha$ may have values as high as $10^3 \text{ cm}^{-1}$ for very high imperfection densities, but in general is considerably less. Absorption of photons by free carriers, cause a transition to higher energy states within the same band or to higher bands. This process can occur over a wide range of photon energies. It involves the absorption of both photons and phonons since both energy and $k$ must be changed in the transition. There is also an optical absorption due to free carriers acting collectively as a kind of electron gas, which is known as plasma resonance absorption.

### 2.1.16.3 Absorption edge

The transition of electrons between the valence and conduction bands in the polymers start at the absorption edge which refer to the minimum or threshold energy at which the absorption coefficient started. The transition is called direct if it is extreme occur due to direct absorption of the incident photonic energy, or can be indirect transition in case of phonon assisted initiation. Mostly all the materials have discriminated absorption edge and the energy gap $E_g$, which related to their chemical properties, hence such criteria can be utilized as a fingerprint in materials characterization (Tauc, 1974).
The absorption constant $\alpha(\nu)$ within the range of $\sim 1 \text{ cm}^{-1}$ up to $10^4 \text{ cm}^{-1}$, is described by the following equation (2.21)

$$\alpha(\nu) = \alpha_0 \exp \left( \frac{h \nu}{E_e} \right)$$  \hspace{1cm} (2.21)

Where, the energy $(E_e)$ is the absorption edge in eV.

2.1.16.4 Optical band gap

At high enough absorption levels ($\alpha > 10^4 \text{ cm}^{-1}$), the absorption constant $\alpha(\nu)$ commonly take place as frequency dependence from which the band gap energy between the conductive band CB and the valence band VB can be deduced, depending on Mott and Davis (1979) which is applicable for UV-spectrum. It correlates between energy band gap $E_g$ and the absorption coefficient $\alpha(\nu)h\nu$ of the composites as in the following equation (2.22)

$$\alpha(\nu)h\nu = B(h\nu - E_{g})^{m}$$  \hspace{1cm} (2.22)

Where $h\nu$ is the energy of the incidence photon, $h$ is the Planck constant, $E_g$ is the optical band gap energy, $B$ is a constant known as the disorder parameter which is dependent on composite composition and independent to photon energy. Parameter $m$ is the power coefficient with the value that determined by the type of possible electronic transitions, such as direct allowed and indirect allowed (Tauc, 1974). The direct transition is the transition of electron from the band $i$ via $k$ band to the final conduction band $f$ by the transition-dipole moments and the electric fields at the surface, while the indirect transition the incident photon is absorbed in the substrate followed by scattering of a photo-excited electron trapped into the intermediate state $k$ band then transfer to final band $f$, as being illustrated in Figure (2.7). All the electrons of the valence band can connect to the empty states of the conduction band by
indirect transition process; however phonons, could also participate in the transition process (Pankove, 1971).

![Diagram](image)

**Figure 2.7** Schematically illustrates the sequence of direct electronic transitions from the initial state $i$ to the final state $f$, or by an indirect process in which the intermediate state $k$ is populated by scattering and relaxation of “hot” electrons, which are photo-excited in the substrate.

2.1.16. 5 UV-Visible absorption spectrophotoscopy

The following Figure (1.8) explains the UV-visible spectroscopy principle, in which a beam of light from a UV-visible light source is separated into its component wavelengths by a prism or diffraction grating. Each monochromatic (single wavelength) beam in turn is split into two equal intensity beams by a half-mirrored device. One beam for the sample (colored magenta), passes through a small transparent container (cuvette) as a sample chamber. The other beam, for reference (colored blue), passes through an identical cuvette. The intensities of these light beams are then measured by electronic detectors and compared. The intensity of the
reference beam, with no light absorption, is defined as \( I_0 \) and the intensity of the sample beam is defined as \( I \). Over a short period of time, the spectrometer automatically scans all the component wavelengths in the manner described. The UV region scanned is normally from 200 to 400 nm, and the visible portion is from 400 to 800 nm (Whiffen, 1971).

If the sample compound does not absorb light of a given wavelength then \( I = I_0 \). However, if the sample compound absorbs light then \( I \) is less than \( I_0 \), and this difference may be plotted on a graph versus wavelength, as shown on the right. Absorption may be presented as transmittance \( (T = I/I_0) \) or absorbance \( (A = \log I_0/I) \). If no absorption has occurred, \( T = 1.0 \) and \( A = 0 \). Most spectrometers display absorbance on the vertical axis, and the commonly observed range is from 0 (100% transmittance) to 2 (1% transmittance). The wavelength of maximum absorbance is a characteristic value, designated as \( \lambda_{\text{max}} \).

Different compounds have very different absorbance. The reference absorption intensity for the system is based on a completely transparent standard compound (non-absorbing). The most commonly used compounds are water, ethanol, hexane and cyclohexane. The compounds with double or triple bonds and heavy atoms are generally avoided, because the absorbance of a sample will be proportional to its molar concentration in the sample cuvette.
Figure 2.8  Schemes for UV-visible spectroscopy principle and steps of taking the spectra.
2.2 previous studies

The following part deals with the previous studies concerning the utilization of PVA hybridized with silver nitrate (AgNO$_3$), in the field of radiotherapy as radiation detector and dosimeter, in these view considerable studies have been introduced:

Karthikeyan, (2005) has been prepared Ag–Polyvinyl alcohol (Ag–PVA) films, and the influence of annealing time upon the variation of cluster size has been analyzed using optical absorption, emission and Fourier-transform infrared (FTIR) spectra. Measurements of optical spectra show that the surface Plasmon resonance lays around 420 nm, and confirm the growth of Ag clusters. Measurements of FTIR spectra were carried out to identify the role of chemical interface damping which influenced the broadening of the absorption band. The emission peak was observed at 540 nm for clusters which were annealed for 1 min as well as 2 min.

The optical absorption spectra of Ag–PVA films annealed at 120 °C at different time duration as shown in Fig. (2.9). All the spectra show the presence of a peak around 420 nm; this peak is characteristic of silver’s SPR. While increasing the annealing time duration the peak intensity also increases, and correspondingly the FWHM becomes lesser. This clearly indicates that while increasing the time duration the mean size of Ag cluster formed also increases. At the same time, a red shift of the peak maximum and a noticeable broadening have been observed for the 10, 15 and 30 min annealed films.
Aleksandra et al, (2007) were prepared the Ag-PVA nanocomposites with different contents of inorganic phase by reduction of Ag\(^+\) ions in aqueous PVA solution by gamma irradiation followed by solvent evaporation. Optical properties of the colloidal solutions and the nanocomposite films were investigated using UV–vis spectroscopy, and the result showed an absorption peak at 420 nm corresponding to PVA/Ag film after irradiation with three different particle size, as shown in Fig. (2.10):
Structural characterization of the Ag nanoparticles was performed by SEM and XRD. Interaction of the Ag nanoparticles with polymer matrix and the heat resistance of the nanocomposites were followed by IR spectroscopy and differential scanning calorimetry (DSC) analysis. IR spectra indicated that Ag nanofiller interact with PVA chain over OH groups. The changes of heat resistance upon the increase of the content of inorganic phase are correlated to the adsorption of polymer chains on the surface of Ag nanoparticles.

Zidan, (1999) prepared Polyvinyl alcohol films, with various AgNO₃ filler mass fractions (≤ 5%), The structural and morphological variations, due to filling and UV-irradiation, were investigated using differential scanning calorimetry (DSC), UV/VIS optical absorption spectroscopy, X-ray diffraction and scanning electron microscopy (SEM). Two different crystalline phases (one is due to the PVA matrix and the other
is attributed to the PVA/Ag+ chelates) were detected besides the PVA amorphous phase, for the non-irradiated and the UV-irradiated (for 2, 4 and 6 h) films as shown in Figures (2.11), (2.4) and (2.12).

Figure 2.11 the UV/VIS optical absorption spectra after IT 2 h for film of AgNO3 filled with PVA system.

Figure 2.12 the UV/VIS optical absorption spectra after IT 4 h for film of AgNO3 filled with PVA system.
The PVA–Ag+ chelates disappeared at 2 h UV-irradiation. It is implied that the structural morphology changes vastly due to the changes in filling level and/or UV-irradiation time. The observed morphological patterns were discussed.

Other study carried out by Kron et al, (1996), in which they said: that if LiF TLD chips of three different thicknesses (0.230, 0.099 and 0.038 g.cm-2) are exposed in the same radiation field it is possible to extrapolate their dose reading to an infinitesimal thin detector size. They used this TLD extrapolation technique to determine the entrance and exit surface dose on a solid water phantom in various photon beams of radiotherapy treatment units. Both build-up and build-down effect could be assessed in dose measurements for telecaesium and megavoltage X-ray beams from a medical linear accelerator while a rapid dose fall-off was observed at the phantom surface in a 120 kVp beam due to electron contamination from the lead glass treatment cone. One of the major advantages of the TLD extrapolation
technique is that the dose can be interpolated and extrapolated for all vital structures within the first millimeter of the skin.

Mutahir et al, (2011), stated the International Commission of Radiological Units (ICRU) sets a tolerance of ±5 % on dose delivery, with more recent data limiting the overall tolerances to ±3 %. One of the best methods for accurate dose delivery and quality check is in vivo dosimetry, while radiotherapy is performed. The present study was carried out to test the applicability of diodes for performing in vivo entrance dose measurements in external photon beam radiotherapy for pelvic tumors and its implementation as quality assurance tool in radiotherapy. During November 2007 to December 2009, in 300 patients who received pelvic radiotherapy on a multileaf-collimator-assisted linear accelerator, the central axis dose was measured by in vivo dosimetry by p-Si diodes. Entrance dose measurements were taken by diodes and were compared with the prescribed dose. Totally 1000 calculations were performed. The mean and standard deviation between measured and prescribed dose was 1.26 ± 2.8 %. In 938 measurements (93.8 %), the deviation was <5 % (1.36 ± 2.9%); in 62 measurements (6.2 %) the mean deviation was >5 % (5.51 ± 2.3 %). Larger variations were seen in lateral and oblique fields more than anteroposterior fields. For larger deviations, patients and diode positional errors were found to be the common factors alone or in combination with other factors. After additional corrections, repeated measurements were achieved within tolerance levels. This study showed that diode-detector-based in vivo dosimetry was simple, cost–effective, provides quick results and can serve as a useful quality assurance tool in radiotherapy. The data acquired in the present study can be used for evaluating output calibration of therapy machine, precision of calculations, effectiveness of treatment plan and patient setup.
Farhat et al, (2011) were measured the entrance and exit dose for patient treated for head and neck tumors. The target absorbed dose was determined from the exit and entrance dose measurement. Twenty patients were evaluated. The results were compared to the calculated values and the midline dose was determinate and compared with the prescribed dose. 80 entrance doses and 80 exit doses measurements were performed. The average difference from expected values was 1.93% for entrance dose (SD 1.92%) and 0.34% for exit dose (SD 4.1%). The target absorbed dose differed from prescribed dose values by 2.94% (1.97%) for the results using the Noël method and 3.34% (SD: 2.29%) with the Rizzotti method. The total uncertainty budget in the measurement of the absorbed entrance and exit dose with diode, including diode reading, correction factors and diode calibration coefficient, is determined as 3.02%. Simple in vivo dose measurements are an additional safeguard against major setup errors and calculation or transcription errors that were missed during pre-treatment chart check.

Tung et al, (2004) they carried out the following measurements using Diode and thermo-luminescence dosimeter (TLD) for linear accelerators of 6 MV photon beams. ESTRO in vivo dosimetry protocols were followed in the determination of midline doses from measurements of entrance and exit doses. Of the fields monitored by diodes, the maximum absolute deviation of measured midline doses from planned target doses was 8%, with the mean value and the standard deviation of -1.0and 2.7%. If planned target doses were calculated using radiological water equivalent thicknesses rather than patient geometric thicknesses, the maximum absolute deviation dropped to 4%, with the mean and the standard deviation of 0.7 and 1.8%. For in vivo dosimetry monitored by TLDs, the shift in mean dose remained small but the statistical precision became poor.
Most relevant study was introduced by Mohammed et al, (2011) they said that for the wider interested applications of polymer hybridized metal nanoparticles composites, poly vinyl-alcohol /silver nanoparticles composites in a form of a film was prepared by in situ irradiation doping technique up to 50 kGy. The effect of radiation upon the composites resulted in reducing the silver ions into black metallic silver, so the general film color changed from white to golden-yellow then black color at 50 kGy as shown in Figures (2.13) and (2.14):

![Color change of PANI nanoparticles polymerized by radiation doping at different doses in PVA blend for 28.6%-AniHCl monomer](image)

Figure 2.14 shows the color change of PANI nanoparticles polymerized by radiation doping at different doses in PVA blend for 28.6%-AniHCl monomer. The picture of the un-irradiated film was taken on a white background.
Figure 2.15 shows the colour change to golden yellow of irradiated PVA/AgNO₃ composites due to reduction of AgNO₃ to Ag⁺ nanoparticles induced by γ-rays. The picture of the un-irradiated film was taken on a white background.

The UV-visible spectroscopy revealed an absorbance band peaking at 425 nm which was increased exponentially with dose increment. The study of UV-spectrogram revealed that the maximum absorbance $A_{\text{max}}$ increased following the particles radius. Scanning electron Microscopy (SEM) revealed shiny nanoparticles of silver cored in Polyvinyl-alcohol PVA with homogeneous distribution and having an average size of 30 nm as well as the XRD spectrum that shows cubic center face of silver nanoparticles in the film and a crystalline peak for PVA reduced by radiation to amorphous phase. Gamma Radiation Synthesis and Characterization of Polyvinyl Alcohol/ Silver Nanocomposites Film.

Shaheen et al, (2013) they studied the investigation of film dosimeters made from polyvinyl alcohol (PVA) films dyed with methyl orange (MO) to enable their use in high dose radiation processing applications was studied. The dosimetric change in
these films at pre and post irradiation was studied spectrophotometrically. Radiolytic bleaching was observed in PVA aided films exposed with Cs137 $\gamma$-source in dose range of 100-200kGy. The effects of pH, dye concentration and film thickness on the radiation response of the film dosimeters were discussed. The stability of MO-PVA films before and after exposure of radiation was also examined at ambient temperature and was found to be higher for long times at pre and post irradiation stages.

They found that the optical absorption spectra of unirradiated and irradiated films were measured in the wavelength range of 300-800nm using a UV/VIS spectrophotometer. The absorption spectra of these unirradiated films show a main absorption band in the visible region peaking at 488nm ($\lambda$ max) for MO-PVA as shown in the Fig (2.15).

![Absorption Spectra](image-url)

Figure 2.16 shows absorption spectra of unirradiated and irradiated MO-PVA films to different absorbed doses.
Mohammad et al, (2013) they studied the effects of gamma irradiation on absorption properties of Carbol Fuchsin (CF) with crystal violet dyed Polyvinyl Alcohol (PVA) blended for possible use in dosimetry and measurement of radiation dose in gamma rays have been studied using UV-Visible spectrophotometer method. The films were irradiated to accumulated doses from (10 to 70) Gy using Co60 gamma ray source at a constant dose rate. The absorption spectra were measured using UV-Visible Spectrophotometer in the wave length range (200 to 900) nm, resulting in a decrease of the absorbance at 560 nm band peak with increasing dose. Regarding to the IR transmittance it’s appearing that for the samples with crystal violet the band at 1716 divided two small bands which become obvious for the irradiated one which indicate also the effect of gamma doses. In this study, the less Full Width at Half Maximum (FWHM), the better is the dosimeter. The FWHM due to sharpness in the band peak and this is useful for determine the fine dosimeter.

In this study, the poly vinyl alcohol based films containing carbolfuchsin and crystal violet dyes after irradiation with gamma ray that radiation induced bletching of these films can be used for dosimetry when this films ex- posed to accumulated doses from (10 to 70 Gy) in wave length range (200 to 900) showed an absorption band at 560 nm. The fitting of the dose-response curves y = ax + b resulted in a linear relationship between dose and net absorbance change. The full width at half maximum (FWHM) in two films which decreased with increasing doses due to sharpness in the band peak. The FWHM can be used to determine any films exposed to high doses and it is used for determining the better dosimeter. Smaller FWHM value resolution enables the system to more clearly separate the peaks within a spectrum.

Ramnani et al, (2007) they have been synthesized Silver clusters on SiO2 support using Co60 gamma radiation, the irradiation of Ag+ in aqueous suspension of SiO2 in the presence of 0.2 moldm_3 isopropanol resulted in the formation of yellow
suspension. The absorption spectrum showed a band at 408nm corresponding to typical characteristic surface plasmon resonance of Ag nanoparticles. The effect of Ag+ concentration on the formation of Ag cluster indicated that the size of Ag clusters vary with Ag+ concentration, which was varied from 4-10-4 to 5-10-3 moldm-3. The results showed that Ag clusters are stable in the pH range of 2–9 and start agglomerating in the alkaline region at pH above 9. The effect of radiation dose rate and ratio of Ag+/SiO$_2$ on the formation of Ag clusters have also been investigated. The prepared clusters have been characterized by X-ray diffraction (XRD) and transmission electron microscopy (TEM), which showed the particle size of Ag clusters to be in the range of 10–20 nm, as shown in Figure (2.16).

![Absorbance vs Wavelength nm](image)

**Figure 2.17** shows the gamma radiation – induced formation of Ag on SiO$_2$ at various radiation doses: (a) 0 kGy, (b) 0.12 kGy, (c) 0.24 kGy, (d) 0.36kGy, (e) 0.54 kGy, (f) 0.66 kGy, and (g) 0.78 kGy.

Zidan, (2002), prepared PVA films with various filling levels of CrF$_3$ and MnCl$_2$. ESR and UV/VIS optical analysis were used to shed more light on the structural modification that occur due to filling with different levels and/or UVirradiation. The
ESR analysis revealed that the spin configuration of CrF$_3$, MnCl$_2$, and CoBr$_2$-filled PVA are different. The filling level dependence of ESR parameters was discussed. The UV-VIS spectral analysis for pure PVA shows absorption bands at 265 and 280 nm, which were assigned to the presence of carbonyl groups. The addition of CrF$_3$ led to the appearance of another bands at 418 and 596 nm. The filling level and/or UV irradiation have no effect on the position of absorption bands but the intensity of these bands has been changed. The addition of MnCl$_2$ led to a new band at about 350 nm due to charge transfer transition. The ligand field parameters and optical energy gaps can be calculated and discussed. The results of optical and ESR analysis indicated that the Cr$^3+$ or Mn$^2+$ are present in its octahedral symmetrical form within the PVA Matrix. SEM micrographs of CrF$_3$ filled PVA is discussed.

Ultraviolet-Visible (UV/VIS) absorption spectra of unirradiated and irradiated samples were carried out in the wavelength range from 200–900 nm as shown in Fig. (2.17) and (2.18).
Figure 2.18 Shows the UV/VIS absorption spectra for films of CrF\textsubscript{3} filled PVA system before UV-irradiation.

Figure 2.19 shows the UV/VIS absorption spectra for films of CrF\textsubscript{3} filled PVA system after UV-irradiation.
CHAPTER THREE
Chapter three
Methodology

3.1 Tools and equipment

3.1.1 Polyvinyl alcohol (PVA)

PVA is one of the important polymer binder and available in the form of powders, fibers and films. It can be obtained from poly (vinyl acetate) (PVAc) by esterification and has distinct crystallinity. The polymer has intermolecular chin of 2.5 Å and consists of 1,3 glycol linkages, in which all hydroxyl groups are arranged along the same side of the chain (Sakurada, 1985). These in turn account for the mechanical strength and strong interactions between different chains. The unit cell of PVA consists of two monomer units of vinyl alcohol (CH$_2$CHOH) (Bunn, 1948) as in Figure 3.1.

![Chemical structure of poly(vinyl alcohol) (PVA) monomer.](image)

PVA is glassy and does not dissolve in water at room temperature unless the ester groups of PVAc are cleaved to a hydroxyl content of 87-89%. It dissolves easily at temperature above glass-transition temperature, $T_\text{g}$ ranging from 60 to 90 °C. It is non-ionic vinyl polymer, tough with film forming capacity due to hydrogen bonding (Ravve, 2000), fibres and tubes forming capability and highly resistant to hydrocarbon solvents (Molyneux, 1983). It is highly resistive to electrical conductivity with low dielectric loss. Its electrical conductivity and charge storage capability can be significantly influence by doping with suitable impurities (Nagaraja, et al., 2002). Due to biocompatibility, PVA has been used in medical devises, materials for drug delivery system, carrier for cell signaling, sizing, adhesives, emulsification and bio-separation membranes (Yano, et al., 2003).
3.1.2 Silver Nitrate (AgNO₃):

Silver nitrate is a chemical compound with a chemical formula AgNO₃ shown in Figure 3.2. This nitrate of silver is the light sensitive ingredient in photographic film and is a corrosive compound. Soluble silver salts tend to be very toxic to bacteria and other lower life species. The compound notably stains the skin giving a blackening colour which is made visible after exposure to sunlight. Silver nitrate is one of the significant compounds in the field of industries due to its potential characteristics such as wider response to electromagnetic radiations i.e. optical properties in addition to electronic, magnetic and catalysis (Wang and Toshima, 1997). It has been used in wider applications as conductive ink, thick film pastes, adhesive for electronic compounds (Lin and Wang, 1996) and photonic and photographic applications (Jin et al., 2001). The characteristics of silver nitrate are has a molecular weight of 169.87, boiling point 444 °C, melting point 212 °C as crystal structure rhombic, decomposed by heat to give Ag, NO₂ and O₂.

![The chemical structure of the silver nitrate compound.](image)

Figure 3.2 The chemical structure of the silver nitrate compound.
3.1.3 $^{60}$Co teletherapy unit:

3.1.4 The source

The $^{60}$Co source is produced by irradiating ordinary stable $^{59}$Co with neutron in a reactor. The nuclear reaction can be represented by $^{59}$Co (n,γ) $^{60}$Co.

The $^{60}$Co source, usually in the form of a solid cylinder disc or pallets, is contained inside stainless capsule and sealed by welding. The double welded seal is necessary to prevent any leakage of the radioactive material. The $^{60}$Co source decay to $^{60}$Ni with the emission of β particles ($E_{\text{max}} = 0.32$ Mev) and two photons per disintegration of energies 1.17 and 1.33 Mev, these two photons constitute the useful treatment beam. The β particles are absorbed in the cobalt metal and the stainless-steel capsules resulting in the emission of bremstrung x-ray and small amount of characteristic x-ray.

A typical teletherapy $^{60}$Co source is a cylinder of diameter ranging from 1.0 to 2.0 cm and is positioned in the cobalt unit with its circular end facing the patient. The fact that the radiation source is not a point source complicates the beam geometry and gives rise to what is known as the geometric penumbra.

3.1.5 Source Housing

The housing for the source is called the source head (Fig. 3.3). It consists of a steel shell filled with lead for shielding purposes and a device for bringing the source in front of an opening in the head from which the useful beam emerges. Also, a heavy metal alloy sleeve is provided to form an additional primary shield when the source is in the off position. A number of methods have been developed for moving the source from the off position to the on position. These methods have been discussed in detail by Johns and Cunningham(10). It will suffice here to mention briefly four different mechanisms: (a) the source mounted on a rotating wheel inside the source head to carry the source from the off position to the on position; (b) the source mounted on a heavy metal drawer plus its ability to slide horizontally through a hole running through the source head-in the ON position the source faces the aperture for the treatment beam and in the OFF position the source moves to its shielded location and a light source mounted on the same drawer occupies the on position of the source; (c) mercury is allowed to flow into the space immediately below the source to shut off the
beam; and (d) the source is fixed in front of the aperture and the beam can be turned on and off by a shutter consisting of heavy metal jaws. All of the above mechanisms incorporate a safety feature in which the source is returned automatically to the off position in case of a power failure.

Figure 3.3 shown the cobalt-60 machine that used to irradiate the films

3.1.6 Beam Collimation
A collimator system is designed to vary the size and shape of the beam to meet the individual treatment requirements. The simplest form of a continuously adjustable diaphragm consists of two pairs of heavy metal blocks. Each pair can be moved independently to obtain a square or a rectangle-shaped field. Some collimators are multivane type, i.e., multiple blocks to control the size of the beam.

3.1.7 UV-Visible spectroscopy:

The UV visible spectroscopy version M350 Double Beam Scanning Spectrophotometer and model (Camspec M350) used in this research as shown in Figure (3.4), has a High performance, double beam optical system, Ultra stable, simultaneous reference subtraction, Range: 190 to 1100nm, and ability to move filter, lamp change points Comprehensive scanning, concentration and kinetics modes and Multi-component analysis - up to 8 components
3.1.8 Optical densitometer:
The optical densitometer is a simple device that shines whitelight onto one side of the developed film and electronically measures the amount oflight reaching the other side. The optical densitometer has a small sensitive area (aperture), typically about 3 mm in diameter, and measurements of OD correspond to that specific area of the film. If the intensity of the light measured with no film in the optical densitometer is given by $I_0$, and the intensity measured at a particular location on a film is given by $I$, then the transmittance ($T$) of the film at that location and the OD are defined as follows:

$$T = \frac{I}{I_0}$$ (3.1)

$$OD = \log_{10}(T) = \log_{10}(\frac{1}{T}) = \log_{10}(\frac{I_0}{I})$$ (3.2)

If the transmission through the film is $T = 0.1 = 10^{-1}$, then $OD = 1$; if the transmission is $T = 0.01 = 10^{-2}$, then $OD = 2$. The relationship between OD and $T$ is analogous to that between pH and the hydrogen ion concentration ($pH = -10\log[H^+])$. The
inverse relationship is $T = 10^{OD}$. The optical densitometer that we used in this work is from PTW model densix as shown in Figure (3.5).

![Optical densitometer](image)

Figure 3.5 shown the optical densitometer that we used in this work from PTW (densix)

### 3.1.9 TLD reader:

The thermoluminescent dosimeter, or TLD, is a type of radiation dosimeter. A TLD measures ionizing radiation exposure by measuring the intensity of visible light emitted from a crystal in the detector when the crystal is heated. The intensity of light emitted is dependent upon the radiation exposure. Materials exhibiting thermoluminescence in response to ionizing radiation include but are not limited to calcium fluoride, lithium fluoride, calcium sulfate, lithium borate, calcium borate, potassium bromide and feldspar.

The two most common types of TLDs are calcium fluoride and lithium fluoride, with one or more impurities to produce trap states for energetic electrons. The former is used to record gamma exposure, the latter for gamma and neutron exposure (indirectly, using the Li-6 (n,alpha) nuclear reaction; for this reason, LiF dosimeters may be enriched in lithium-6 to enhance this effect or enriched in lithium-7 to reduce
As the radiation interacts with the crystal it causes electrons in the crystal's atoms to jump to higher energy states, where they stay trapped due to intentionally introduced impurities (usually manganese or magnesium) in the crystal, until heated. Heating the crystal causes the electrons to drop back to their ground state, releasing a photon of energy equal to the energy difference between the trap state and the ground state. The electrons can also drop back to ground state after a long period of time; this effect is called fading and is dependent on the incident radiation energy and intrinsic properties of the TLD material. As a result, each material possesses a limited shelf life after which dosimetric information can no longer be obtained. This varies from several weeks in calcium fluoride to up to two years.

It can be used both for environmental monitoring and for staff personnel in facilities involving radiation exposure, among other applications. The TLD reader sued in this research was PCL3 from PTW. The PCL3 is an automatic reader to measure accumulated dose signals from thermos luminescence dosimeters. Up to 80 detectors can be evaluated in one cycle. The detectors are stored in a magazine, automatically processed and loaded into a second magazine. TL chips of 4.5 mm diameter and 0.8 mm thickness and TL powder are preferably used, alternatively TL rods and TL films are suitable. The detectors are heated by contact. The heating temperature of maximum 600 °C is controlled automatically with an accuracy of better than 1%. The measuring range covers seven decades with a linear response of up to 12 Gy using standard TLD material and up to 500 Gy using material LiF:Mg,Ti. The background reading is corrected automatically before starting each measurement. An internal reference light provides for automatic calibration of the photomultiplier. The dose reading can be calibrated in the radiation units Gy, Sv, rem or rad or in the unit C (electrical charge). The PCL3 system is prepared for TLD evaluation under nitrogen atmosphere. The modular cabinet design of the device presents very easy access to the mechanical and electronic components for maintenance and repair. A computer software for control and evaluation is included in the standard package.
3.2 Method:

3.2.1 Preparations of PVA/AgNo3 composite sample films

The PVA stock was supplied by SIGMA (Mw = 72,000 g/mol, 99 – 100% hydrolyzed). The PVA solutions were made by dissolving 3.00 g PVA powder in 100 ml distilled water at temperature of 80 °C on a beaker as shown in Figure (3.7). The solution was magnetically stirred throughout at that temperature for 3 hours and then left to cool at room temperature.
After cooling to room temperature, we took the PVA solution into dark room and hybridized with 0.2 wt% of AgNo3 and stirred for 2h. Then the PVA /AgNo3 solution poured in a petri-dish to form films by casting method in dark room. Then the PVA/AgNo3 blend solution was divided into two, 40 ml each(The remaining 20 ml evaporated due to heat) and spread into a specially made glass caster, 10 cm in diameter and left to dry at ambient temperature in dark room for at least 3 days to remove water. Then films will be peeled off the petri-dish, cut into small films 2x2cm, loaded in sealed dark plastic packs (dental film envelope).

3.2.2 Irradiation of film samples

All film samples together with TLD were irradiated with doses in range of 0 – 15 Gy, using $^{60}$Co $\gamma$-rays irradiation facility (MDS nordion, equinox), from radiation and Isotopes centre Khartoum as shown in Figure (3.8).
Figure 3.8 shown the films position during irradiation by $^{60}$Co teletherapy machine.

The samples stick at the Perspex phantom on the entrance and exit area of the radiation beam; the diameter of the Perspex phantom is 40x40cm with 10cm thickness, the films were embedded at right angle to direction of the radiation beam as shown in Figure (3.8).

The dose rate on the exact date of measurement July 27, 2012 was 1.35 Gy/min. The new dose rate was calculated based on the half-life, $T_{1/2}$, of the source, i.e. the elapsed time in which the activity of the source reduced by half, is equal to 5.27 years by the equation (3.3)

$$
\dot{D} = \dot{D}_0 \exp\left(-\ln 2 \frac{t}{T_{1/2}}\right)
$$

(3.3)

Where $\dot{D}_0$ is the dose rate at time $t = 0$. In May 2005, the dose rate of the source was calibrated using Ferrous Sulphate dosimeter. Fricke dosimeter was then irradiated in the calibrated beam to the doses that the solution is well characterized. The dosimeter was read using UV-Vis spectrophotometer to measure the absorbance of FeSO$_4$ at the wavelength of 304 nm. The deviation of the current value to standard value is of main interest for chamber stability and is maintained to within ± 1%. The deviation was calculated using the following equation (3.4).

$$
Deviation = \frac{2.75 \times 10^2 \times \Delta A}{1 + 0.007(T - 25^\circ C)}
$$

(3.4)
where $\Delta A$ is the changing absorbance, $T$ is the temperature in Celsius. Response of Fricke dosimeter was compared to the secondary standard chamber and the consistencies of by these methods were maintained to be less than $\pm 3\%$. The solution was then sent for standardization using gamma chamber and also for calibration of routine dosimeter. The routine check source measurements were performed every three months in order to assess the long term stability of the reference standard chamber.
CHAPTER FOUR
Chapter four

Results

The following chapter deals with highlighting of prepared PVA\AgNO₃ composites films and their relevant characterization after irradiation with doses in the range of radiotherapy together with utilization in radiation dose measurement with comparison with TLD.

4.1 Color formation change

Figure 4.1 shows the change in the measured parameters for the studied samples due to amount of doses.

4.2 UV-visible spectrometer

Figure 4.2 shows the UV-spectrum for PVA\AgNO₃ film receiving γ-radiation dose 1-15 Gy as entrance applied dose.
Figure 4.3 shows the UV-spectrum for PVA\AgNO₃ film receiving γ-radiation dose 1-15 Gy as exit dose.

Figure 4.4 show the correlation between the applied γ-radiation doses 1-15 Gy and the absorption coefficient at entrance and exit doses.
4.3 Optical density:

Figure 4.5 shows the correlation between dose in Gy and the relevant optical density in (a. u.).
4.4 TLD results:

Figure 4.6 showed the correlation between the applied doses (given dose) in Gy and TLD reading in Gy for entrance and exit dose.

Figure 4.7 shows the correlation between TLD Dose reading in Gy and the relevant optical density in (a. u).
Figure 4.8 shows the correlation between the absorption in Gy and the attenuation coefficient $\mu$
CHAPTER FIVE
Chapter five

Discussion, Conclusion and recommendations

5.1 Discussion

The following part is deal with result discussion as appear in chapter four:

Figure (4.1) shows the colour change of the irradiated films receiving different radiation doses (1, 2, 4… 12 and 15 Gy). The color changed from white (un-irradiated film) then to light yellow, golden then to brown and dark brown following the increment of radiation dose. Such color change has been reported by (Vladimir et al, 2012) and it was due to reduction of Ag⁺ by the formed reducing species as hydrated electron (eaq), hydrogen atom radical (H°) and hydroxyl radical (OH°). Same study has been reported by Ramnani et al, (2007) and Mohammed et al, (2011, 2013).

Also figure 4.2 shows the UV-spectrum for PVA\AgNO₃ films after irradiation with γ-radiation receiving same doses in figure (4.1). It showed that the irradiated films having absorption band peaking at the wavelengths 200 and 410 nm which are related to criteria of the pure PVA and PVA/AgNO₃ respectively. The absorption bands increased following the radiation dose increment, such phenomena has been stated by Mohammed et al and Ramnani et al. The absorptions in figure (4.2) is related to the applied doses at the entrance of the radiation field, therefore they showed more prominent peaks than in figure (4.3) which is related to the absorption at exit site of radiation beam.

The figure 4.4 shows the correlation between the applied γ-radiation doses 1-15 Gy and the absorption coefficient at \(\lambda =200\) and \(\lambda =410\) nm for both entrance and exit doses. It showed that the absorption coefficient is increase linearly with increasing of applied dose based in the following equation \(y = 0.09x + 0.62\) and \(y = 0.1x + 0.1\).
for entrance and exit dose respectively, where \( y \) refers to the absorption in (au) and \( x \) refers to the dose in Gray (Gy) with a correlation coefficient \( r^2 = 0.99 \), indicating the sensitivity of the composite film to radiation which in turn showed the possibility of using such types of film as radiation detector and personal dosimeter after suitable build in an electronic circuit.

Figure 4.5 shows the correlation between applied dose in Gy and the relevant optical density in a u, for entrance and exit doses in the range of 1-15 Gy at a peak of 200 and 410nm. It shows that there is a linear proportional correlation between the two parameters (dose in Gy and optical density) in the absorption band of 200 and 410 nm, however the relation of absorption in the range of 410 nm is strong as: \( r^2 = 0.98 \), for entrance and exit doses ; which ascribed to the sensitivity of the silver to radiation and further more encouraging the application of PVA/Ag film as a radiation detector or monitoring based on optical density change. The correlation could be fitted to the equation of the following form: \( y = 0.04x + 0.12 \) and \( y = 0.03x + 0.99 \) for entrance and exit doses respectively, where \( y \) refers to optical density and \( x \) refers to the applied dose in Gy.

The figure 4.6 shows showed the correlation between the applied doses (given dose) in Gy and TLD reading in Gy for entrance and exit dose. It has been noticed that: the average entrance dose was 7.06± STD 4.7 Gy while the exit dose was 3.98± STD 2.8Gy in average i.e. the average absorption between entrance and the exit was 3.08 ± STD 1.9 Gy, such result confirming the accuracy of TLDs in measurement of the doses at entrance and exit of the phantom.

The figure 4.7 shows the correlation between TLDs dose reading in Gy and the relevant optical density a.u for entrance and exit doses in the range of 1.1-14.26 Gy at a peak of 200 and 410 nm. It shows that there is a linear proportional correlation between the two parameters in the absorption band of 200 and 410 nm however the relation of absorption in the range of 410nm is strong as \( R^2 =0.987 \) for both entrance and exit dose which ascribed to the
sensitivity of the silver to radiation and further more encouraging the application of PVA/Ag film as a radiation detector or monitoring based on optical density change. And the correlation could be fitted to the equation of the following form: $y = 0.04x + 0.117$ and $y= 0.031+ 0.099$ for entrance and exit dose respectively, where $y$ refers to optical density and $x$ refers to the applied dose in Gy. Mohammed et al, (2013) and Mohammed et al, (2011). Such equation (entrance dose) could be utilized to take the synthesized films as personal dosimeters in x-ray clinics at rural areas where the TLD reader is not available, depending on the optical density change.

In figure 4.8 the correlation between the dose absorption in Gy and the attenuation coefficient $\mu$ was appear. It has been deduced that: the attenuation coefficient of the phantom determined based on the exponential law, which is equal to 0.1 in average, such result is agreed with the manufacture of tissue equivalent materials which is equal to 0.14, Hill et al, (2008). And the relationship could be fitted in the following exponential equation: $y = 0.083e^{-0.09x}$, where $y$ refers to Attenuation coefficient $\mu$ and $x$ refers to radiation absorption in Gy.
5.2 conclusions

According to the analyzed results of the following experimental, the researcher could conclude that:

- The PVA\Ag composites films have been synthesized using controlled temperature and casting method.
- The irradiation of the films with gradation induces obvious color change from white (un-irradiated film) then to light yellow, golden then to brown and dark brown following the increment of radiation dose.
- The irradiated films showed absorption band peaking at the wavelengths 200 and 410 nm and the intensity of peaks in proportional increment with radiation dose.
- The absorption coefficient was increased in linear form following the increment of of applied dose based in the following equation
  \[ y = 0.09x + 0.62 \] for entrance and exit dose respectively, where \( y \) refers to the absorption in (au) and \( x \) refers to the dose in Gray (Gy). The correlation coefficient \( r^2 = 0.99 \) indicating the sensitivity of the composite film to radiation which in turn showed the possibility of using such types of film as radiation detectors or personal dosimeter.
- There is strong correlation between the applied dose and the optical density as \( r^2 = 0.96 \), and the equation could be fitted in the following equations:
  \[ y = 0.04x + 0.12 \] and \( y = 0.03x + 0.99 \) for entrance and exit doses respectively, where \( y \) refers to optical density and \( x \) refers to the applied dose in Gy.
- The average entrance dose was 7.06± STD 4.7 Gy while the exit dose was 3.98± STD 2.8 Gy in average, and the average absorption between entrance and the exit was 3.08 ± STD 1.9 Gy.
- There is linear proportional correlation between the applied dose and optical density which could be fitted in the following equation: \( y = 0.04x + 0.117 \) and \( y = 0.031 + 0.099 \) for entrance and exit dose respectively, where \( y \) refers to optical density and \( x \) refers to the applied dose in Gy.

- Also there is applicable relationship \((R^2 = 0.63)\) between the dose and the attenuation coefficient \( \mu \) of phantom based to the following equation: \( y = 0.083e^{-0.09x} \), where \( y \) refers to Attenuation coefficient \( \mu \) and \( x \) refers to radiation absorption in Gy.
5.2 Recommendations

This research could consider an initial step for measuring radiation in radiotherapy department, accordingly we recommend by the following recommendations:

- PVA/Ag composite films can be used as a local tool for in vivo dosimeter system to measure the entrance and exit dose in radiotherapy centers in Sudan because of its good result, fast, easy and cheap assessment tool.
- Further work is needed to support this research especially for measuring absorbed dose at deference depths in radiotherapy using PVA/Ag composite film.
- We recommend using modern techniques to synthesize the PVA/Ag film with suitable envelope.
- The color change of the films could be utilized as light filters and the surface Plasmon resonance could be utilized as local in vivo irradiation therapy as well as in signal magnification in case of Raman scattering.
- Radiation therapy phantom could be developed from such composites (PVA\Ag) for dose histogram, mapping, and simulation.
- Further work is needed to know how the film thickness is affecting by the radiation dose.
References


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http://www.camspec.co.uk/m350.htm.

http://www.ptw.de/pcl3_automatic_tld_reader.html.

ICRU (1976), Determination of absorbed dose in a patient irradiated by beams of x or gamma rays in radiotherapy procedures. ICRU report 24, Bethesda, Maryland.


Appendix A

TLD reading for entrance and exit dose

<table>
<thead>
<tr>
<th>G.D in Gy</th>
<th>TLD En in Gy</th>
<th>TLD Ex in Gy</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>1.1</td>
<td>0.403</td>
</tr>
<tr>
<td>2</td>
<td>1.98</td>
<td>0.966</td>
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<tr>
<td>4</td>
<td>3.91</td>
<td>1.98</td>
</tr>
<tr>
<td>6</td>
<td>5.90</td>
<td>3.29</td>
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<tr>
<td>8</td>
<td>7.79</td>
<td>4.55</td>
</tr>
<tr>
<td>10</td>
<td>9.69</td>
<td>5.60</td>
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<tr>
<td>12</td>
<td>11.81</td>
<td>6.67</td>
</tr>
<tr>
<td>15</td>
<td>14.26</td>
<td>8.35</td>
</tr>
</tbody>
</table>
## Appendix B

Applied dose for each film and optical density for entrance and exit dose

<table>
<thead>
<tr>
<th>Applied Dose</th>
<th>Entrance (OD)</th>
<th>Exit (OD)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1Gy</td>
<td>0.17</td>
<td>0.14</td>
</tr>
<tr>
<td>2Gy</td>
<td>0.19</td>
<td>0.15</td>
</tr>
<tr>
<td>4Gy</td>
<td>0.24</td>
<td>0.19</td>
</tr>
<tr>
<td>6Gy</td>
<td>0.26</td>
<td>0.22</td>
</tr>
<tr>
<td>8Gy</td>
<td>0.33</td>
<td>0.26</td>
</tr>
<tr>
<td>10Gy</td>
<td>0.35</td>
<td>0.30</td>
</tr>
<tr>
<td>12Gy</td>
<td>0.41</td>
<td>0.32</td>
</tr>
<tr>
<td>15Gy</td>
<td>0.44</td>
<td>0.34</td>
</tr>
</tbody>
</table>
Appendix C

The correlation between the applied $\gamma$-radiation doses 1-15 Gy and the absorption coefficient at entrance and exit doses

<table>
<thead>
<tr>
<th>Applied dose in Gy</th>
<th>Entrance absorption coefficient</th>
<th>exit absorption coefficient</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>0.73</td>
<td>0.134</td>
</tr>
<tr>
<td>2</td>
<td>0.86</td>
<td>0.31</td>
</tr>
<tr>
<td>4</td>
<td>0.98</td>
<td>0.569</td>
</tr>
<tr>
<td>6</td>
<td>1.20</td>
<td>0.853</td>
</tr>
<tr>
<td>8</td>
<td>1.42</td>
<td>1.02</td>
</tr>
<tr>
<td>10</td>
<td>1.64</td>
<td>1.24</td>
</tr>
<tr>
<td>12</td>
<td>1.76</td>
<td>1.43</td>
</tr>
<tr>
<td>15</td>
<td>2.15</td>
<td>1.71</td>
</tr>
</tbody>
</table>