Comparative analysis of antioxidant and phenolic content of chloroform extract/fraction of *Terminalia chebula*

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**ABSTRACT**

In the present study, two chloroform extracts of fruits of *Terminalia chebula* viz. “CHL1” and “CHL 2” prepared by maceration and sequential method respectively was compared for their antioxidant efficacy and phenolic content. The extraction procedure of plant material plays an important role in the activity of phytochemicals. Also, the assessment of antioxidant capacity of phytochemicals cannot be executed precisely by any single method due to complex nature of phytochemicals as multiple reaction characteristics and mechanisms can be involved. So, no single assay could accurately reflect comparison in a mixed or complex system. Therefore in the present study the comparison of extracts was done by using most widely used assays viz. DPPH, deoxyribose, reducing power, chelating power and lipid peroxidation assay. Furthermore, the UV-Vis spectrum of both extracts and the correlation between total phenolic content was examined in order to give an orientation to the search of phytochemicals responsible for their activity. From the results, it was concluded that antioxidant activity and phenolic compounds were predominant in the ‘CHL 2’ prepared by sequential method. The present study enlightening the useful extraction procedure of plant material.

**INTRODUCTION**

Redox regulation is an imperative process for homeostasis in human physiology and presently scientific studies are focused to comprehend its detailed mechanism. Free radical originates in the body by a myriad of biochemical processes that leads to oxidative stress. It is commonly accepted that, in a situation of oxidative stress, more reactive oxygen species (ROS) and reactive nitrogen species (RNS) are generated. Oxidative stress, the shift of the redox balance through oxidative potentials may damage biological molecules altering cell function and leads to various diseases such as neurodegenerative disorders, cancer, cardiovascular diseases, atherosclerosis, cataracts, and inflammation [1-3]. Antioxidants, molecules that inhibit or prevent oxidation of a substrate, evolved to protect biological systems against damage induced by ROS/RNS [4, 5]. A sophisticated, co-operative array of antioxidant defense mechanisms called antioxidant network is found in biological systems. But in stress conditions the defense system of the body fails to provide protection against there harmful radicals, so the antioxidants are provide externally [6]. The health promoting effect of antioxidants from plants is thought to arise from their potential effects on the reactive oxygen/nitrogen species [7].
Antioxidant properties elicited by plant species have a full range of prospective applications in human healthcare. The history of the use of medicinal plants for alleviation of diseases has its origin in primitive times. It has been recognized that there is an inverse association between consumption of some fruits and vegetables and degenerative diseases, which could be partly attributed to their antioxidant activity [8-10]. Many medicinal plants are explored for their pharmacological activities but, indubitably the plant kingdom still holds several species of plants containing substances of medicinal value that have yet to be discovered. It is reported that phenolic compounds in plants possess strong antioxidant activity and may help to protect cells against the oxidative damage caused by free radicals [11,12]. Due to the presence of the conjugated ring structures and hydroxyl groups, many phenolic compounds have the potential to function as antioxidants by scavenging superoxide anion [13], singlet oxygen [14], and lipid peroxy radicals [15], and stabilizing free radicals.

As part of our efforts to find antioxidants from edible fruits, we have investigated the antioxidant activity of fruits of *Terminalia chebula*, which has been used extensively in the Ayurveda to treat various ailments. It is found in the sub Himalayan tracks from Ravi to West Bengal, Assam and in all deciduous forests of India, specifically in Madhya Pradesh, Bihar, Assam and Maharashtra. The fruits contain about 30% astringent substances like chebulinic acid, tannic acid, gallic acid and chebulagic acid [16-19]. From fruit kernels, palmitic, stearic, oleic, linoleic, arachidic and behenic acids are also isolated. The fruits are mild laxative, stomachic, tonic, alterative, adaptogen, hepatoprotective, febrifuge, antispasmodic, expectorant, anti-asthmatic, antiviral and hypoglycaemic [20-23]. It is also useful in ophthalmia, hemorrhoids, dental caries, bleeding gums, ulcerated oral cavity and in many other diseases. This characteristic makes it a good therapeutic prospect of study.

**MATERIALS AND METHODS**

**EXTRACTION PROCEDURE**

The fruits of the *T. chebula* were purchased locally from market, washed, dried and finally ground to fine powder. The chloroform extract was prepared by maceration and sequential methods.

**Maceration method**

1 kg of dried fruit powder of *T. chebula* was washed with hexane to remove extraneous matter. After drying of fruit powder the chloroform was added and the mixture was stirred for 24 hours. The suspended solid was filtered through Whatman no.1 filter paper and the filtrate was collected. This procedure was performed thrice to get three filtrates of chloroform and residue. Chloroform filtrate were combined and dried at room temperature in petri plates. The green coloured gummy solid left behind was named as ‘CHL 1’ for all further applications (Flow chart 1).

**Sequential method**

1 kg of fruit powder of *T. chebula* was suspended in methanol and the mixture was kept for 24 hrs at room temperature. The residue was filtered through Whatman no.1 filter paper and filtrate was collected. The filtrate was dried at room temperature by putting in petri plates to get methanol extract. 100g of methanol extract was dissolved in 20% aqueous methanol and put into the separatory funnel. This layer was washed with hexane to remove fatty compounds. Then the chloroform was added in separatory funnel and mixed properly. After some time the two layers were formed which were separated to get chloroform filtrate and 20% aqueous methanol stock. The three filtrates of chloroform were collected and dried at room temperature in petri plates to get chloroform extract named as ‘CHL 2’ (Flow chart 2).

**CHEMICAL ANALYSIS**

**Determination of total phenolic content**

The total phenolic content of the extract was determined using Folin-Ciocalteu method [24]. To 100μl of extract was added 900μl of water. To this 500μl of FC reagent was added. This was followed by the addition of 1.5ml of 20% Sodium carbonate. The mixture was shaken thoroughly and allowed to stand for 2 hours. The volume of mixture was made up to 10ml with distilled water and absorbance was observed at 765nm. The phenolic content was calculated as gallic acid (mg/g) equivalents.

**Spectroscopic analysis of extracts**

The ‘CHL 1’ and ‘CHL 2’ extracts of *T. chebula* were analyzed by UV spectroscopy. For this the
solution of ‘CHL 1’ and ‘CHL 2’ was prepared in spectroscopic grade methanol in the concentration of 1mg/10ml, diluted four times and a spectrum was recorded on UV-Visible spectrophotometer (Shimadzu-1601).

**ANTIOXIDANT TESTING**

**DPPH assay**

The Hydrogen donor activity of the ‘CHL 1’ and ‘CHL 2’ extracts was measured by 1, 1 diphenyl - 2 - picrylhydrazyl (DPPH) method [25]. The reaction mixture contained 200μl of different extracts concentrations and 2 ml of DPPH. The reaction mixture was observed at 517nm against a blank, which did not contain extract. The % age inhibition was calculated as:

\[
\text{% Inhibition} = \frac{B_0-B_1}{B_0} \times 100
\]

where, \(B_0\) is the absorbance of control, \(B_1\) is the absorbance of reaction mixture.

**Deoxyribose assay**

This was done by non-site specific and site-specific methods with modifications [26]. Stock solution of EDTA (1mM), FeCl₃ (10mM), ascorbic acid (1mM), H₂O₂ (10mM) and deoxyribose (10mM) were prepared in distilled water. In non-site specific assay, 0.1ml of EDTA, 0.01ml of FeCl₃, 0.1 ml of H₂O₂, 0.36 ml of deoxyribose, 1ml of extracts concentrations (10-100 μg/ml), 0.33 ml of phosphate buffer (50mM, pH 7.4) and 0.1 ml of ascorbic acid were added in sequence. The mixture was incubated at 37°C for 1 hour. 1ml of the incubated mixture was mixed with 1 ml of 10% trichloroacetic acid and 1 ml of thiobarbituric acid (0.025M NaOH) and heated for one hour on water bath at 80°C and pink chromogen developed, which was measured at 532 nm. In site-specific assay EDTA was replaced with phosphate buffer. The % age inhibition was calculated as:

\[
\text{% Inhibition} = \frac{B_0-B_1}{B_0} \times 100
\]
Flow chart 2: Chloroform extract of fruits of *T.chebula* prepared by sequential method.

where, $B_0$ is the absorbance of control, $B_1$ is the absorbance of reaction mixture

**Reducing power assay**

The reducing activity of the ‘CHL 1’ and ‘CHL 2’ extracts was determined according to the method of Oyaizu [27]. 1ml of extract/fractions (50-300μg/ml) was prepared in distilled water and mixed with 2.5 ml of phosphate buffer (0.2M, pH 6.6) and 2.5 ml of 1% potassium ferricyanide. The mixture was incubated at 50°C for 20 min. 2.5 ml of 10% trichloacetic acid was then added to the mixture and centrifuged at 3000 rpm for 10 min. 1ml of aliquot of supernatant was mixed with 2.5ml of distilled water and 0.5 ml of FeCl₃ (0.1%) and absorbance was measured at 700nm. Increase in absorbance was interpreted as increased reducing activity.
Comparative analysis of antioxidant and phenolic content of chloroform

Chelating power assay
The chelating activity of extracts was measured as given by Dinis et al. [28] with little modifications. 1 ml of extract with different concentrations was mixed with 3.5 ml of methanol, and then the mixture was mixed with ferrous chloride (2 mM, 0.1 ml) and ferrozine (1 mM, 0.2 ml) for 10 min at room temperature. The absorbance was measured at 562 nm against a blank in which the extract was not added. The % age inhibition was calculated as:

\[
\% \text{ Inhibition} = \frac{B_0 - B_1}{B_0} \times 100
\]

where, \(B_0\) is the absorbance of control, \(B_1\) is the absorbance of reaction mixture

Lipid peroxidation assay
This method was done according to the method of Halliwell and Guttridge (29). Normal albino rats of the Wistar strain were used for the preparation of liver homogenate. The perfused liver was isolated, and 10% (w/v) homogenate was prepared using a homogenizer at 0-4°C with 0.15 M KCl. The homogenate was centrifuged at 3000 rpm for 15 min, and clear cell-free supernatant was used for the study of in vitro lipid peroxidation. Different concentrations of extracts mixed with 1 ml of 0.15 M KCl and 0.5 ml of rat liver homogenates were added to the test tubes. Peroxidation was initiated by adding 100 μl of 0.2 mM ferric chloride. After incubation at 37°C for 30 min, the reaction was stopped by adding 2 ml of ice-cold HCl (0.25 N) containing 15% trichloroacetic acid (TCA), 0.38% thiobarbituric acid (TBA) and 0.5% butylated hydroxytoluene (BHT). The reaction mixtures were heated at 80°C for 60 min. The samples were cooled and centrifuged, and the absorbance of the supernatants was measured at 532 nm. An identical experiment was performed to determine the amount of lipid peroxidation obtained in the presence of inducing agents without any extract. The % age inhibition was calculated as:

\[
\% \text{ Inhibition} = \frac{B_0 - B_1}{B_0} \times 100
\]

where, \(B_0\) is the absorbance of control, \(B_1\) is the absorbance of reaction mixture

STATISTICAL ANALYSIS
The data presented as Mean ±SE of three independent experiments

RESULTS AND DISCUSSION
In the present study the comparison of two chloroform extracts was done prepared by different extraction method. From the analysis of total phenolic content, it was observed that the chloroform extract prepared by maceration method has low phenolic content as compared to chloroform extract prepared by sequential method. The phenolic content of ‘CHL 1’ extract was 0.9 mg/g of GAE, while it was 3.15 mg/g of GAE in ‘CHL 2’ extract. To further confirm the presence of phenolic content, the UV analysis of ‘CHL 1’ and ‘CHL 2’ extract was done which again confirmed the presence of phenolic compounds. The UV spectrum of the ‘CHL 1’ extract gives absorbance at \(\lambda_{max}\) in region 260 nm with end absorption around 200 nm (Figure 1). The small structureless absorption in region 260 nm

Figure 1: The UV-visible spectra of ‘CHL 1’ acquired by maceration method.
indicates the absence of polyphenolic and phenolic compounds and indicates the presence of aliphatic/terpenoids in this. The UV spectrum of ‘CHL 2’ extract revealed the absorption around \( \lambda_{\text{max}} = 362 \text{nm} \), which indicated the presence of polyphenolic compounds (Figure 2). Phenolic compounds exhibited two major absorption bands in the ultraviolet/visible region i.e. a first band in the range between 320 and 380 nm and a second band in the 250 to 285 nm range [30]. So, it can be concluded that the ‘CHL 2’ extract contain phenolic compounds as foremost phytochemical.

The H-donor ability of ‘CHL 1’ and ‘CHL 2’ extracts was measured and compared by using DPPH assay. In this assay, at 500\( \mu \)g/ml dose ‘CHL 1’ extract exhibited 8.1% inhibition and ‘CHL 2’ extract exhibited 58.2 % inhibition (Figure 3). The reaction time of ‘CHL 1’ extract was 30-35 minutes and for ‘CHL 2’ extract 15-20 minutes. Possible mechanism of DPPH scavenging was suggested to be through reduction (protonation) of this radical by antioxidant compound to a more stable DPPHH form [31]. Because of its unpaired electron, DPPH has an absorption maximum at 520 nm and as it gets reduced as electron paired off in the presence of a radical scavenger. The absorbance decreases stoichiometrically with respect to the number of electrons taken up. So, from the results it can be alleged that ‘CHL 2’ has the hydrogen donating ability which may be due to phenolic content. The labile hydrogen donating capacity of ‘CHL 2’ extract was assessed during this study can be linked to important mechanism of phenolic antioxidant action [32]. The data of the present investigation suggests that the extracts are able to donate 

Figure 2: The UV-visible spectra of ‘CHL 2’ acquired by sequential method.

![Figure 2](image1)

Figure 3: Effect of different concentrations of ‘CHL 1’ and ‘CHL 2’ extract of T.chebula in DPPH assay.

![Figure 3](image2)
electrons and therefore should be able to stabilize the reactive radical to unreactive species. In the absence of the extract, no decrease in DPPH and absorbance was seen with time however, it started decreasing in the presence of extract.

The hydroxyl radical scavenging activity of ‘CHL 1’ and ‘CHL 2’ was compared by using non-site specific and site specific deoxyribose assay. In non-site specific the ‘CHL 1’ extract showed 36.2% and ‘CHL 2’ extract showed 57.5 % inhibition at 300μg/ml of dose (Figure 4). In case of site-specific the ‘CHL 1’ extract showed 27.5% and ‘CHL 2’ extract showed 65.3 % inhibition at same dose (Figure 5). In this method the deoxyribose is degraded by OH radical formed from Fenton reaction and form malonaldehyde (MDA) fragments, which in turn react with thiobarbituric acid (TBA) to produce a TBA reactive chromophore that was detectable at 532nm [33,34]. Hence, the deoxyribose assay has become a useful experimental tool for investigating the ability to react with OH radicals. Our results suggested that ‘CHL 2’ extract is not only effective scavengers of OH in this system but also iron chelator, as it showed more activity in site-specific assay than in non-site specific assay. Thus the compounds, which are able to chelate iron preferentially and change the metal complex in a less redox active form will protect deoxyribose against damage. From the UV-spectroscopy, it is clear that the ‘CHL 2’ extract showed maximum absorbance in polyphenolic region which can be fore most cause of its activity.
The reducing capacity of a compound may serve as a significant indicator of its potential antioxidant activity [35, 36]. Literature reports have shown that the reducing power of bioactive compounds is associated with antioxidant activity [37-40]. In this assay, the antioxidant capacity is measured on the basis of the ability to reduce ferric (III) ions to ferrous (II) ions in the presence of ‘CHL 1’ and ‘CHL 2’ extracts. The ‘CHL 1’ extract showed 0.255 absorbance and ‘CHL 2’ showed 0.555 absorbance at 500 μg/ml concentration (Figure 6). Less activity of ‘CHL 1’ extract may be due to the absence of phenolic compounds or co-occurrence of other compounds like fatty acids, which hindered its antioxidative activity and made these moieties non-functional. The ‘CHL 2’ extract showed considerable increase in absorption, which may be due to the presence of phenolic compounds as confirmed by UV spectroscopy. Difference in activity of chloroform extracts may be due to variations in extraction procedure which influenced extractability and stability of polyphenolic compounds.

Transition metals contain one or more unpaired electrons that enable them to participate in one-electron transfer reactions so they are powerful catalysts of autoxidation reactions like conversion of H₂O₂ to -OH in the Fenton reaction and in the decomposition of alkyl peroxides to the highly reactive alkoxyl and hydroxyl radicals [41, 42]. The chelating property of the phytochemicals can contribute significantly towards antioxidant behavior. In the present study it was observed that the iron chelating effect of ‘CHL 1’ extract was 7.5 % and ‘CHL 2’ extract was 57.3% at 250 μg/ml of concentration (Figure 7).
It was clear that ‘CHL 2’ extract interfered with the formation of ferrous-ferrozine complex, suggesting that it have marked iron chelating activities and capture ferrous ion before the formation of ferrozine.

In order to compare the chloroform extracts are capable of reducing in vitro oxidative stress, the lipid peroxidation assay that determines the production of malondialdehyde and related lipid peroxides in living system was carried out. Likewise in earlier experiments, here also the ‘CHL 2’ extract indicated the highest ability of 62.3% and ‘CHL 1’ extract possessed 17.3% inhibition at highest tested dose (Figure 8). Polyphenols in ‘CHL 2’ extract can combine with hydroxyl radical and convert them into water molecules and stabilize them to inhibit MDA production or they chelate the active iron sites so that the Fenton reaction could not take place. So, it may be supposed that different polyphenols are responsible for antioxidative potential of ‘CHL 2’ extract.

CONCLUSION

In the end, it can be concluded that ‘CHL 2’ extract of T. chebula prepared by sequential method exhibited good hydrogen donating, radical scavenging, metal chelating, reducing and antioxidant potential. The results obtained from this study showed that extraction method plays an important role in antioxidant potential of any extract. Also, employment of more than one test method specific to a radical species gives a better estimate of comparative antioxidant potential of a test compound.

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