Al-hijamah and oral honey for treating thalassemia, conditions of iron overload, and hyperferremia: toward improving the therapeutic outcomes

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Abstract: Iron overload causes iron deposition and accumulation in the liver, heart, skin, and other tissues resulting in serious tissue damages. Significant blood clearance from iron and ferritin using wet cupping therapy (WCT) has been reported. WCT is an excretory form of treatment that needs more research efforts. WCT is an available, safe, simple, economic, and time-saving outpatient modality of treatment that has no serious side effects. There are no serious limitations or precautions to discontinue WCT. Interestingly, WCT has solid scientific and medical bases (Taibah mechanism) that explain its effectiveness in treating many disease conditions differing in etiology and pathogenesis. WCT utilizes an excretory physiological principle (pressure-dependent excretion) that resembles excretion through renal glomerular filtration and abscess evacuation. WCT exhibits a percutaneous excretory function that clears blood (through fenestrated skin capillaries) and interstitial fluids from pathologocial substances without adding a metabolic or detoxification burden on the liver and the kidneys. Interestingly, WCT was reported to decrease serum ferritin (circulating iron stores) significantly by about 22.25% in healthy subjects (in one session) and to decrease serum iron significantly to the level of causing iron deficiency (in multiple sessions). WCT was reported to clear blood significantly of triglycerides, low-density lipoprotein (LDL) cholesterol, total cholesterol, uric acid, inflammatory mediators, and immunoglobulin antibodies (rheumatoid factor). Moreover, WCT was reported to enhance the natural immunity, potentiate pharmacological treatments, and to treat many different disease conditions. There are two distinct methods of WCT: traditional WCT and Al-hijamah (WCT of prophetic medicine). Both start and end with skin sterilization. In traditional WCT, there are two steps, skin scarification followed by suction using plastic cups (double S technique); Al-hijamah is a three-step procedure that includes skin suction using cups, scarification (shartat mihjam in Arabic), and second skin suction (triple S technique). Al-hijamah is a more comprehensive technique and does better than traditional WCT, as Al-hijamah includes two pressure-dependent filtration steps versus one step in traditional WCT. Whenever blood plasma is to be cleared of an excess pathological substance, Al-hijamah is indicated. We will discuss here some reported hematological and therapeutic benefits of Al-hijamah, its medical bases, methodologies, precautions, side effects, contraindications, quantitative evaluation, malpractice, combination with oral honey treatment, and to what extent it may be helpful when treating thalassemia and other conditions of iron overload and hyperferremia.

Keywords: Al-hijamah, prophetic medicine, cupping therapy, phlebotomy, iron chelation therapy, oral honey

Introduction

Cupping therapy is the term given to treatment modalities using sucking cups that are applied to the skin surface at different anatomical points. It includes both dry cupping
therapy (DCT) and wet cupping therapy (WCT). WCT is an excretory form of treatment that needs more research as a safe, economic, time-saving technique without serious side effects.1 WCT does not require sophisticated equipment, expensive disposables, or extensive effort to practice.1 There are no serious limitations, precautions, or side effects to discontinue WCT. Interestingly, WCT has solid scientific and medical bases that explain its effectiveness in treating many disease conditions differing in etiology and pathogenesis according to the evidence-based Taibah mechanism (Taibah theory)1 (Supplementary video 1). WCT includes both the traditional WCT (skin scarification and suction method), and the more comprehensive WCT of prophetic medicine, Al-hijamah (skin suction, scarification, and suction method). We will discuss here some reported hematological and therapeutic benefits of WCT, its medical basis, methodologies, WCT-induced reduction in serum iron and ferritin, combination with oral honey, and to what extent this may be helpful when treating thalassemia, hemochromatosis, sideroblastic anemia, and other conditions of iron overload and hyperferremia.

**Important reported therapeutic and hematological benefits of WCT**

A single session of WCT (blood cupping therapy, Al-hijamah) was reported to decrease serum ferritin (circulating iron stores) significantly by about 22.25% in healthy subjects (from 129±59.2 to 100.6±45.8 ng/mL).2 The same session of WCT cleared blood significantly of triglycerides, low-density lipoprotein (LDL) cholesterol, total cholesterol, and uric acid.2 In rheumatoid arthritis, WCT cleared patients’ blood of immunoglobulin antibodies (rheumatoid factor) in a time-dependent manner,3 which may increase the therapeutic benefits of WCT. In another report, frequent WCT was reported to decrease serum iron, causing iron deficiency that brought about iron deficiency anemia.4

Potentiation of the natural immunity in the form of physiological leukocytosis and an increased number of natural killer cells were reported after applying WCT.5 That was confirmed by the study of Zhang et al, who reported that traditional WCT decreased serum levels of the immune inhibitory cytokines (eg, immunoglobulin E [IgE]), interleukin-4 (IL-4), IL-10) and decreased the number of T cytotoxic cells (CD8+) cells. In the same subjects, WCT increased the production of endogenous immunostimulatory cytokines (eg, interferon-gamma, IL-2, complement-3, complement-4, immunoglobulins [IgA, IgG, and IgM]) and activated cells of the immune system (eg, T helper cells), causing an increased ratio of T helper/T cytotoxic cells,6 which may help in treating hepatitis viral infections that may complicate frequent blood transfusions.

Being a mechanical modality of treatment, WCT has no chemical or pharmacological antagonism with pharmacological treatments such as iron chelation therapies. As mentioned before, WCT was reported to clear blood significantly of excess pathological substances, which may facilitate and potentiate the related pharmacological and therapeutic effects of coadministered pharmacotherapy.1 Moreover, WCT was reported to significantly decrease the blood pressure, and improve the lipid profile and treat different diseases, such as hypertension and hyperlipidemia.1,2 WCT was reported to exert pharmacological potentiation when treating different diseases and to alleviate drug-induced side effects;1,3 for example, WCT-induced leukocytosis counteracted methotrexate-induced leucopenia when treating rheumatoid patients with Al-hijamah combined with methotrexate.13 Interestingly, skin wounding (eg, the minor skin scarifications done during WCT) was reported to enhance the production of endogenous nitric oxide through upregulating the expression of nitric oxide synthase (NOS).1,6 Neuronal NOS messenger ribonucleic acid (mRNA) is upregulated and neuronal NOS protein expression peaks at the late stages of skin wound healing.7 Moreover, inducible NOS mRNA was reported to increase by about tenfold,8 peak 24 hours after wounding, and persist for a few days.9 Nitric oxide exerts many important roles, including antioxidant effects,10 vasodilator functions, and antimicrobial effects,11 among others.1 Recently, Al-hijamah was reported to exert multiple therapeutic health benefits; for example, nonspecific biochemical clearance of serum and interstitial fluids, excretory benefits, antinociceptive (analgesic) effects, anti-inflammatory effects, hemodynamic effects, antiviral effects, anticancer effects, antiallergic effects, respiratory benefits, and neurological health benefits. Moreover, Al-hijamah was reported to exert nonspecific-tissue protective benefits, cardioprotective effects, hepatoprotective benefits, nephroprotective effects, and neuroprotective effects, among others. Therefore, the reported antihypertensive and antihyperlipidemic effects of Al-hijamah may be beneficial in guarding against the development and progression of premature atherosclerosis that is encountered in some diseases, such as thalassemia. We reported Al-hijamah as a promising suggested treatment for hematological conditions characterized by iron overload; eg, hemochromatosis, thalassemia, sideroblastic anemia, and others.11

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Indices for evaluating therapeutic effects of Al-hijamah

Recently, we reported the novel therapeutic indices of Al-hijamah for evaluating the health benefits gained from practicing Al-hijamah for treating different disease conditions.\(^1\) Such indices included excretion index, purification index, immunological potentiation index, pharmacological potentiation index, and clinical therapeutic index (CTI).

Excretion index (excretion value, EV) is the quantity of noxious substances excreted after the end of the session of Al-hijamah in quantity units/volume units. EV can be calculated by determining the difference in levels of noxious substances before and after Al-hijamah (in quantity units/volume units). An example for that was the decrease in serum ferritin from 129.4 ng/mL (before Al-hijamah) to 100.6 ng/mL (after Al-hijamah):

\[
EV = (129.4 - 100.6) = 28.8 \text{ ng/mL.}
\]

Plasma clearance index, or purification index (PI), is the percentage of purification of plasma from any noxious substance or component estimated at different time points after Al-hijamah. PI can be calculated by estimating the difference in the concentration of a noxious substance before and after Al-hijamah divided by its initial plasma concentration level. Using the abovementioned example, PI for plasma from serum ferritin equals

\[
100 \times \left( \frac{129.4 - 100.6}{129.4} \right) = 22.25\%.
\]

This means that Al-hijamah induced clearance of more than one-fifth of initial serum ferritin in a single session, which may be promising when treating conditions of iron overload.

Pharmacological potentiation index measures the degree of pharmacological potentiation gained when conventional pharmacological treatments are administered concomitantly with practicing Al-hijamah. It also compares the gained therapeutic benefits versus using conventional pharmacological treatment as a sole treatment. When treating rheumatoid arthritis patients using conventional pharmacological treatment, the initial serum rheumatoid factor was 129.75 to 131.47 IU/mL, which decreased to 122 IU/mL when conventional pharmacological treatment was used as a sole treatment.\(^3\) A combination of conventional pharmacological treatment plus Al-hijamah caused serum rheumatoid factor to decrease to 51.46 IU/mL. Pharmacological potentiation index for decreasing serum rheumatoid factor:

\[
\left( \frac{122 - 51.46}{122} \right) = 60.2%.
\]

Thus, combining Al-hijamah with pharmacological treatment increased the gained therapeutic effects more than ten times.

Immunological index measures the degree of immunological potentiation gained after Al-hijamah; eg, natural killer cell lymphocytosis. It can be calculated by measuring the number of natural killer cells before and after Al-hijamah. A percentage count of natural killer cells before Al-hijamah was 8.5% and increased to 11.33% after Al-hijamah. II for the increase in the count of natural killer cells can be calculated as

\[
\left[ 100 \times \left( \frac{11.33}{8.5} \right) \right] = 133\%.
\]

Thus, natural immunity using natural killer (NK) cells increased by about 133% after Al-hijamah.

CTI is the improvement percentage of an investigated clinical parameter (eg, diastolic blood pressure value, systolic blood pressure value, pain intensity value, disease activity value, and others) after Al-hijamah (as a sole treatment or as a combined treatment) was measured at different time points versus the same value before Al-hijamah. Diastolic blood pressure was 81.1 mmHg before Al-hijamah and significantly decreased to 75 mmHg after Al-hijamah.\(^2\) The improvement in diastolic blood pressure equals

\[
(81.1 - 75) = 6.1 \text{ mmHg.}
\]

When calculating the percentage improvement versus the initial value of systolic blood pressure, CTI can be derived from the formula:

\[
\text{CTI} = \left[ 100 \times \frac{6.1}{81.1} \right] = 7.5\%.
\]

Thus, diastolic blood pressure decreased by 7.5% in healthy subjects after Al-hijamah.

Scientific principles of WCT and Al-hijamah

Making use of simple scientific facts in physiology and histology may improve treatment outcomes. For example, human skin has a superficial network of fenestrated dermal capillaries\(^13\) (Figure 1A) that may help percutaneous pressure-dependent and size-dependent filtration and blood clearance upon application of external suction pressure; WCT utilizes a physiological filtration and an excretory principle for inducing blood clearance. Capillary pores and fenestrations are 6–12 nm (and may reach up to 100 nm in diameter) (Figure 1B),\(^12,14\) which cannot filter intact blood cells (having diameters in microns).\(^14\) This may carry a lot of similarities with renal glomerular filtration. This confirms
the fact that Al-hijamah utilizes a physiological principle for inducing blood clearance (pressure-dependent filtration followed by excretion) that is better than the chemical methods of blood clearance, eg, dialysis (concentration-dependent excretion). All this constituted the basis for Taibah mechanism (Taibah theory) for scientific basis of Al-hijamah and cupping therapy.

Moreover, skin comprises three layers: epidermis, dermis, and hypodermis. Epidermis is the most superficial layer, and does not contain blood capillaries. The most superficial layer of the epidermis is the horny cell layer that is the keratinized epithelium layer (stratum corneum). This keratinized cell layer constitutes the skin barrier (10–20 µm) that provides the function of protecting the skin and the underlying structure through preventing the entry and exit of substances through the skin. Immediately below the stratum corneum, there are the viable epidermal cell layers (50–100 µm thick), which are responsible for generating the stratum corneum.

The dermis (1–2 mm thick) is the second layer of the skin that lies immediately beneath the epidermis. The dermis constitutes the mechanical support for the skin. Unlike the epidermis, the dermis contains fenestrated skin capillaries that are suitable for pressure-dependent filtration upon application of negative suction pressure, as that done during Al-hijamah. Interestingly, the ultrastructure of the skin vasculature confirms that the arterial blood supply of the skin comes through the cutaneous arteries that run in the hypodermis, then give rise to branches that anastomose to form the cutaneous capillary plexus at the dermis–hypodermis junction. That cutaneous capillary plexus gives rise to the subpapillary plexus of fenestrated capillaries that lie at the dermoepidermal junction. The subpapillary plexus gives rise to capillary loops in each dermal papilla. Venous drainage of the skin is arranged in plexuses corresponding to the arterial supply. Based on that, skin scarification done during Al-hijamah should be so superficial as to just open the skin barrier and reach the superficial dermal capillaries.

Scarifying the skin deeply during Al-hijamah is a malpractice that may cause damage to the fenestrated dermal capillaries and impair the filtration function exerted during Al-hijamah, as we will discuss in the side effects and malpractice of Al-hijamah. Huber et al reported that many different methods for creating
suction pressure can be utilized for sucking skin inside sucking cups during cupping therapy. However, some variable differences in pressure may occur and govern the proper choice of the most suitable method to exert such suction pressure. Huber et al studied the mean suction pressure produced during mechanical suction, lighter flame 2 cm suction, lighter flame 4 cm suction, and alcohol flame, and found that large differences in suction magnitude were produced. Negative suction was $-200\pm30$ hecta Pascal (hPa) ($-150\pm23$ mmHg) when using 2 cm flame, $-310\pm30$ hPa ($-232.5\pm23$ mmHg) with 4 cm flame, $-560\pm30$ hPa ($-420\pm23$ mmHg) with burning an alcohol-soaked cotton swab, and $-270\pm16$ hPa ($-202\pm12$ mmHg) with rubber balloon mechanical suction. The pressure produced through the rubber balloon was the easiest technically, moderate in intensity, maximal in safety (not harmful), and was the most reproducible when repeating cupping therapy.17

Methods of practicing WCT

DCT is a one-step therapeutic modality that involves skin suction only and is done mainly for the purpose of pain relief or to stimulate blood flow at a given anatomical area of the human body.11 WCT includes both cupping and bloodletting. There are two reported methodologies of WCT: traditional WCT and Al-hijamah (WCT of prophetic medicine). In both, sterilization using alcohol or povidone iodine in a strict aseptic environment is necessary. Traditional WCT has two steps: skin scarification (puncturing) followed by skin suction (cupping); ie, the double S method or puncturing and cupping method of WCT.16 Al-hijamah has three main steps: skin suction (cupping step using sucking cups) (Figure 2A–C), skin scarification (puncturing skin) (Figure 3A–C), and second skin suction (cupping) (Figure 4A–C); ie, the triple S method or cupping, puncturing, and cupping method of WCT.1,11 The methodology of using traditional WCT for treating many different disease conditions was reported in many published research studies from Germany,19 Korea,20 and the People’s Republic of China.21–24 The methodology of traditional WCT described in those published studies is different from Al-hijamah, which was described in reports from Saudi Arabia25 and other countries.26,27

Each suction step during WCT is a pressure-dependent filtration step; ie, Al-hijamah has two filtration steps versus one filtration step for traditional WCT, which allows also a longer duration of pressure-dependent filtration during Al-hijamah than during traditional WCT.1,11 Based on this, Al-hijamah was reported to provide better filtration, better excretory functions, and less blood loss than traditional WCT.18

Traditional WCT is a partial form of Al-hijamah; ie, all diseases treated by traditional WCT can be treated by Al-hijamah.1,11 Indications for practicing Al-hijamah were previously reported.11 Al-hijamah is indicated whenever an excess noxious substance or fluid is to be removed from plasma and interstitial fluids. Those excess harmful substances may be causative substances for the disease process or pathogenesis; eg, excess uric acid in gout (causing gouty arthritis) and excess iron in thalassemia and iron overload conditions. Both can be referred to as disease-causing substances. When the excess noxious substances are formed during disease pathogenesis; eg, oxidized LDL in thalassemia patients (induced by excess iron), which can be referred to as disease-related substances. Both disease-causing substances and disease-related substances can be referred to as causative pathological substances. Al-hijamah is indicated when the serum level of any causative pathological substances increases in blood or interstitial fluids enough to require clearance.11 Moreover, other therapeutic benefits of Al-hijamah11 may help in treating different disease conditions, such as musculoskeletal pain conditions and neuropsychiatric conditions.11

Skin scarification during Al-hijamah (shartat mihjam in Arabic) (Figure 3A) is a vital step, as it opens the skin barrier and transmits the suction pressure to the fenestrated dermal skin capillaries to favor the excretion of local intercellular fluids and capillary filtration, causing excretion of small-sized particles1,18 (serum iron, serum ferritin, liberated hemoglobin and fragments of hemolyzed cells excluding intact blood cells) from skin capillary blood (Figure 1B). Shartat mihjam refers to the superficial skin scarifications (0.1 mm–0.2 mm in depth) that open the skin barrier to transmit the suction pressure around skin dermal capillaries to facilitate the suction-induced, pressure-dependent filtration process of blood circulating in the fenestrated skin capillaries during Al-hijamah. The superficial nature of skin scarifications ensures that no damage will occur to the fenestrated skin capillaries, which ensures preservation of the pressure-dependent filtration function. To ensure the superficial nature of the skin scarifications, skin incisions done during shartat mihjam should be short (1–2 mm) in length, vertical, in parallel rows, multiple, evenly distributed, confined to the skin uplifting (skin dome), and preceded and followed by a suction step. All that may guarantee the superficial nature of skin scarification and the best conditions for transmitting the negative suction pressure to the fenestrated dermal skin capillaries. Skin scarifications done during traditional WCT cannot be described as shartat mihjam, as those scarifications are not preceded by a suction step.11 Filtered fluids (plasma-like fluids) may come out through shartat mihjam with repeated suctions using sucking cups (Figure 4C). Al-hijamah has been reported to exert a percutaneous nonspecific clearance.
Figure 2 First step of Al-hijamah (first suction step).

Notes: (A) Cups are put mainly at the interscapular regions and upper aspect of the back of trunk (Kahel region over seventh cervical vertebra). (B) Skin upliftings are created and become prominent after removal of cups. (C) Hair follicles become more prominent within the skin upliftings.
Al-hijamah for thalassemia, iron overload, and hyperferremia

Figure 3 Second step of Al-hijamah (skin scarification, shhartat miham in Arabic).
Notes: (A) Skin scarifications should be confined to skin upliftings, superficial (0.1–0.2 mm in depth), short (1–2 mm in length), multiple, and evenly distributed. (B) Cups should be applied immediately after scarifying skin upliftings. (C) Salah’s technique for safe practice of Al-hijamah at special anatomical sites: a simple technique aiming at avoiding possible injury of the underlying anatomical structures (eg, nerves or blood vessels during skin scarification step during Al-hijamah). A fold of the skin uplifting is pinched out between the thumb and index fingers of the left hand to make sure that it is away from the underlying anatomical structures, while the right hand makes superficial scarifications in the pinched out skin fold. Another skin fold is then taken, and so on until finishing the skin uplifting properly.

of blood and interstitial spaces. Increasing the number of Al-hijamah sessions or sucking cups at many anatomical sites may increase blood clearance.

Al-hijamah, also called bloodletting cupping therapy or blood cupping therapy in Arabic countries, is a simple therapeutic modality of WCT that is practiced for treating many diseases. Health professionals practicing Al-hijamah should receive adequate medical education and evaluation. Unfortunately, few research studies have focused on Al-hijamah. Detailed scientific bases of Al-hijamah were...
Figure 4 Third step of Al-hijamah (second suction step).

Notes: (A) Clearance of blood during Al-hijamah. External pressure applied through sucking cups helps filtration of small molecules through fenestrated skin capillaries. Small molecules in iron overload include iron, ferritin, liberated hemoglobin, and debris of hemolyzed or fragmented blood cells. (B) Collected coagulated bloody fluids excreted through Al-hijamah. (C) Filtered fluids (plasma-like) coming out through shartat mihjam with repeated skin suctions.

Abbreviation: RBC, red blood cell.
reported in the evidence-based Taibah theory (Taibah mechanism).\textsuperscript{1,18} Al-hijamah is more comprehensive than the worldwide traditional WCT.\textsuperscript{1,18} Detailed differences between Al-hijamah and traditional WCT were previously reported.\textsuperscript{11,18} In traditional WCT, skin scarifications are done on the surface of the intact skin (not uplifted skin), which may produce pain during the procedure (due to lack of the hypothetic or anaesthetic effect gained during the first suction step). Moreover, lack of the first suction step in traditional WCT (present in Al-hijamah) prevents the formation of a fluid barrier that creates a mechanical barrier between the superficial skin layers and the dermal capillaries. That fluid barrier created during the first step of Al-hijamah separates the fenestrated dermal capillaries from the skin surface (Figure 4A) and protects the dermal capillaries from damage or injury during the superficial skin scarifications. Based on this, the possibility of injuring the fenestrated dermal capillaries using superficial scarifications during traditional WCT is greater than the possibility of injuring dermal capillaries with superficial scarifications during Al-hijamah. It is important to decide which type of WCT (Al-hijamah or traditional WCT) was done when writing or reading research articles about WCT. Some authors may write WCT when Al-hijamah is meant in the methodology section,\textsuperscript{27} which may confuse the reader.

**Proper practice, precautions, contraindications, and side effects of Al-hijamah**

Al-hijamah is a safe therapeutic procedure when practiced properly by a qualified, well-trained, and licensed practitioner. It is rare to face side effects when the criteria for proper practice of Al-hijamah are fulfilled. Strict sterilization is a must in the beginning, throughout the whole procedure, and at the end of Al-hijamah. The most important precaution is to take care regarding selecting the best anatomical sites for cup application.\textsuperscript{28} It is essential to practice Al-hijamah in specialized clinics in hospitals. In addition, moderate suction pressure inside cups (less than \(300 \text{ mmHg}\)) for 5 minutes is the best. It is better if Al-hijamah is not practiced near the course of superficial veins or nerves to guard against injury to those structures. It is not advisable to practice Al-hijamah in patients with high-risk pregnancy, recent burns (inflamed tissues), or attacks of severe bleeding. Circulatory shock, uncontrolled bleeding disorders, uncontrolled coagulation disturbances, conditions of severe anemia, and conditions of active bleeding are relative contraindications for performing Al-hijamah. Once the abovementioned conditions are corrected and the patient’s condition improves, Al-hijamah can be practiced when there is a strong therapeutic indication, taking into account the risk/benefit ratio. When Al-hijamah is highly indicated at some special anatomical sites (near the course of superficial veins or nerves), risk/benefit ratio should be applied and Al-hijamah should be done using Salah’s technique\textsuperscript{28} (Figure 3C) for safe practice of Al-hijamah where a skin fold of the skin uplifting is pinched out between the thumb and index fingers of the left hand to make sure that it is away from the underlying anatomical structures while the right hand makes superficial scarifications in the pinched-out skin fold. The suction step should be manual using a handheld pump\textsuperscript{11} and should not be done using cotton or alcohol ignition to avoid skin burns or fire accidents. Suction time should be about 5 minutes, and suction pressure should be moderate (three to four maximal suction times using the handheld pump).

Shartat mihjam (Figures 3A, B and 4A) is an Arabic term that means superficial skin scarifications (0.1 mm in depth to preserve the structure of the superficial fenestrated dermal capillaries) that are confined to the skin dome (skin uplifting created after the first suction step). Skin scarifications of Al-hijamah denote the hand skills that differentiate one skillful practitioner from another. Skin scarifications should be multiple, longitudinal (1–2 mm in length; ie, not pinpoint pricks), in parallel rows, equally distributed, and productive. Malpractice of Al-hijamah is caused by any violation of those criteria and may carry side effects. Practitioner skill is required to do shartat mihjam properly for better blood clearance benefits that can be evaluated and quantitated using the previously mentioned indices of Al-hijamah. Frequency of performing Al-hijamah varies according to disease condition and response to treatment. It can be done every other week, monthly, twice per year, or yearly. Every case should be individualized according to the patient’s condition and follow-up in light of current treatment.

There are no absolute contraindications for practicing Al-hijamah. Proper practice of Al-hijamah necessitates the presence of an experienced practitioner to perform Al-hijamah and an eligible patient having a disease condition that will benefit from treatment using Al-hijamah. Al-hijamah can be done for both preventive and therapeutic health purposes.\textsuperscript{1,18} Al-hijamah should be done in a sterile, well-equipped environment (hospital or specialized clinic) using the triple S steps (suction, scarification, and suction) of Al-hijamah. In the vast majority of cases, Al-hijamah is done by applying sucking cups on points on the back of the trunk.\textsuperscript{28} This carries the advantages of a simpler technique (the flat surface of
the back is easier for large cup application for better blood clearance) and cosmetic benefit (the back is a hidden area covered by clothes).

The most important points of malpractice causing possible side effects include deep skin scarifications (>0.5 mm in depth) causing damage of the superficial fenestrated capillaries (causing disturbed Al-hijamah-induced filtration function), long skin incisions causing bleeding, large skin incisions causing postcupping scars, strong suction pressure (≥300 mmHg) may cause skin ecchymosis and Taibah sign,11 suction using cotton ignition may burn the skin, prolonged suction time causing skin ecchymosis, and pinpoint pricks causing inefficient clearance function. However, after the acute condition improves, re-evaluation of the patient’s general condition is needed before practicing Al-hijamah where the risk/benefit ratio should be applied. In the literature, side effects of cupping therapy were mainly related to malpractice of traditional DCT and traditional WCT, but not Al-hijamah. As the methodology is similar in all three, reported side effects should be considered and should be divided into common and rare side effects.

Common side effects include skin bruises, Taibah sign11 (due to excessive suction pressure or prolonged suction duration causing dermal capillary rupture), bleeding (due to deep or long skin scarification), keloid scarring, and burns (when using cotton or alcohol ignition). Rare serious side effects may include acquired hemophilia A,29 hemorrhagic stroke,30 factitious panniculitis,31 reversible cardiace hypertrophy, and iron deficiency anemia (due to too frequent sessions of traditional WCT with extensive bleeding caused by an unqualified practitioner).32,33

Pathophysiology of iron overload and hyperferremia
Iron overload is a major health problem in thalassemia, hemochromatosis, sickle cell anemia, sideroblastic anemia, and others that may cause tissue damage (involving the heart, liver, skin, pancreas, and endocrine glands, among others) and serious complications in children and adults. Mean iron level in the skin of thalassemia patients increased by more than 100% of patients. Unfortunately, iron chelation therapy does not treat the abovementioned causes of hemolysis.

Mechanisms of hemolysis in thalassemia
Mechanisms of hemolysis in thalassemia (an example of iron overload) need careful understanding, medical education, and evaluation, being the source of iron (from liberated hemoglobin of hemolyzed cells) in addition to the increased intestinal absorption of iron. Hemolysis in thalassemia is multifactorial in origin; eg, due to precipitation of globin chains in the erythrocytes, erythrocyte membrane skeletal protein defects,16 presence of natural antigalactosyl IgG antibodies on thalassemic erythrocytes,37 alloimmunization (or erythrocyte autoimmunization due to autoantibodies) in transfusion-dependent thalassemic patients,37 sensitivity of thalassemic erythrocytes to oxidative stress (due to increased oxidants as malondialdehyde and decreased antioxidants as polyunsaturated fatty acids),36 and excessive oxidation of the unstable globin chains with subsequent release of oxygen-free radicals.36 Red blood cell destruction by human monocytos may be related to changes in the intracellular iron and ferritin concentrations and phenotype.38

Complications of excessive frequent blood transfusions and excessive hemolysis may aggravate the iron overload condition and may cause blood-borne infections such as viral hepatitis (C or B). Unfortunately, iron chelation therapy does not treat the abovementioned causes of hemolysis.

Iron is high in the interstitial fluids and in the serum of iron overload patients. This may be explained by the fact that interstitial fluid is formed by filtration of capillary blood at the arterial ends of capillaries and is absorbed later at the venous ends of capillaries.1,18 Moreover, the pathophysiology of pulmonary hypertension in beta-thalassemia may be mediated through iron overload, reduced bioavailability of nitric oxide, hemolysis, and hypercoagulopathy.39 As skin iron deposition is high in iron overload, treatment modalities using percutaneous iron excretion may be promising.

Oxidative and pathological roles exerted by excess serum iron and ferritin
Iron is a major oxidant in vivo.40 Iron-induced generation of reactive oxygen species is responsible for the damaging effects induced by iron released by isoferritins (containing iron) on lymphocyte functions.41 Iron overload causes oxidative stress,
Table 1 Reported pathological roles of ferritin in iron overload conditions; eg, thalassemia

<table>
<thead>
<tr>
<th>In thalassemic patients without hepatitis viral infection:</th>
<th>In thalassemic patients with hepatitis viral infection:</th>
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<tbody>
<tr>
<td>High serum ferritin is:</td>
<td>High serum ferritin is:</td>
</tr>
<tr>
<td>• An index of liver iron overload in thalassemia patients (not having viral hepatitis or ascorbic acid deficiency).</td>
<td>• Closely related to the liver iron concentration. Adequate chelation therapy usually protects against liver fibrosis.</td>
</tr>
<tr>
<td>• Associated with immunological suppression.</td>
<td>• Useful in the follow-up of patients receiving long-term transfusional treatment using the ferritin/ALT ratio, especially when acute or chronic liver cell damage may interfere with iron overload by increasing serum ferritin values.</td>
</tr>
<tr>
<td>• Associated with increased growth rates of cancer cells and infectious organisms.</td>
<td>• Associated with hepatitis C virus infection (causing raised AST activity and serum ferritin concentration compared with seronegative patients).</td>
</tr>
<tr>
<td></td>
<td>• Significantly associated with the hepatic fibrosis.</td>
</tr>
<tr>
<td></td>
<td>• A risk factor (with serum triglycerides and total cholesterol) for impaired glucose tolerance and diabetes mellitus.</td>
</tr>
<tr>
<td></td>
<td>• Associated with hepatitis C virus antibody positivity and HCV-RNA by PCR.</td>
</tr>
<tr>
<td></td>
<td>• Associated with high liver transaminases (ALT, AST), anti-HCV seropositivity, raised liver transaminases, hemochromatosis status, and liver fibrosis.</td>
</tr>
<tr>
<td></td>
<td>• Associated with viral hepatitis (genotype 1) with higher rate of splenectomy.</td>
</tr>
<tr>
<td></td>
<td>• Observed in patients with TTV-HCV co-infection compared with patients with TTV infection alone.</td>
</tr>
<tr>
<td></td>
<td>• Accounting in part for the enlargement of hepatoduodenal ligament nodes in thalassemia.</td>
</tr>
<tr>
<td>Low serum ferritin is:</td>
<td>Low serum ferritin is:</td>
</tr>
<tr>
<td>• Associated with sustained virological response to ribavirin.</td>
<td>• Associated with increased growth rates of cancer cells and infectious organisms; eg, cirrhosis and chronic hepatitis. This may cause the formation of antiferritin antibodies and circulating immune complexes.</td>
</tr>
<tr>
<td>• Reported in complete responders to interferon therapy compared with the values for partial and nonresponders before starting therapy.</td>
<td>• Associated with sustained virological response to ribavirin.</td>
</tr>
<tr>
<td>• Observed in patients who responded to interferon-alpha monotherapy.</td>
<td>• Observed in patients who responded to interferon-alpha monotherapy.</td>
</tr>
</tbody>
</table>

**Abbreviations:** ALT, alanine aminotransferase; AST, aspartate aminotransferase; HCV, hepatitis C virus; PCR, polymerase chain reaction; RNA, ribonucleic acid; TTV, Torque teno virus.

---

depletion of endogenous antioxidants, and lipid peroxidative damage. Iron may induce the formation of the major atherogenic factor (oxidized LDL), causing atherosclerosis (Table 1). Serum cholesterol was reported to increase iron deposits that could be identified in the arterial walls in conditions of experimental hypercholesterolemia and in iron-overloaded hypercholesterolemia. Lipoperoxides in liver and spleen homogenates were high in conditions of iron overload.

Moreover, young thalassemic patients showed an atherogenic lipid profile where serum triglycerides, total cholesterol, apolipoprotein A, and carotid intima-media thickness (CIMT) were significantly elevated. Serum high-density lipoprotein (HDL) was significantly lowered. In thalassemic patients, CIMT was positively correlated with age, hemoglobin F (Hb F), ferritin, and cholesterol levels. Interestingly, both serum iron and triglycerides may be involved in the pathogenesis of LDL oxidation, as evidenced by the report that high levels of oxidized LDL antibodies promoted atherosclerosis in patients with beta-thalassemia. That was confirmed by another report in which iron was a potent catalyst of oxidative reactions and macrophage-mediated LDL oxidation in atherogenesis. In addition, released heme may have injurious effects that result from its potent catalysis of LDL oxidation. Moreover, circulating iron stores (serum ferritin) were reported to be associated with vascular damage in patients with nonalcoholic fatty liver disease.

**Iron overload suppresses patients’ immunity**

Iron overload is usually associated with hyperferremia in conditions of beta-thalassemia major, dyserythropoiesis, hereditary hemochromatosis, and chronic diffuse liver diseases; eg, cirrhosis and chronic hepatitis. This may cause the formation of antiferritin antibodies and circulating immune complexes. Increased body stores of iron may affect the immunoregulatory balance, resulting in increased growth rates of cancer cells, infectious organisms, and opportunistic pathogens. Iron overload may complicate the clinical management of pre-existing acute and chronic diseases. Iron and ferritin impaired a variety of immunological functions. Ferritin-associated iron induced neutrophil dysfunction in hemosiderosis. Iron overload, especially with hyperferremia, may exert an immune-suppressing effect through altering the distribution of T lymphocytes in various compartments of the immune system, suppressing the immunological functions of the complement system (classic and alternative pathways), decreasing the T helper (CD4) proliferative capability, decreasing T helper numbers, decreasing T helper activity, increasing the CD8/CD4 ratios, impairing the generation of cytotoxic T cells, enhancing the suppressor T cell numbers and activity, suppressing the antibody-mediated and mitogen-stimulated phagocytosis (by monocytes and macrophages), inducing alterations in the T lymphocyte subsets, inducing modification of lymphocyte distribution in different immunological locations, and altering immunoglobulin secretion.

There is a reported progressive dysfunction of monocytes associated with iron overload in patients with thalassemia major. Moreover, there is an impaired neutrophil defense against *Yersinia enterocolitica* in patients with iron overload who are undergoing dialysis.
Negative correlations between T helper/T cytotoxic ratio and malondialdehyde levels were obtained in the blood and spleen tissues during experimental iron overload, indicating that lipid peroxidation may play a role in the immunological abnormalities observed in experimental hemosiderosis. Secondary hemosiderosis may be accompanied by a decrease in the phagocytic functions of neutrophils, as evidenced by the report that neutrophils from patients with secondary hemosiderosis contain excessive amounts of autotoxic iron and have increased cellular iron and ferritin content. Iron concentrations were four to five times more elevated in secondary hemosiderosis than in healthy subjects. This iron accumulation may be toxic for neutrophils and may explain the three-fold higher risk of bacteremia reported in those patients.

Infections with virulent pathogens are commonly encountered in patients having iron overload; eg, *Listeria monocytogenes* meningitis. Although iron-binding proteins such as transferrin and lactoferrin are bacteriostatic in vitro, iron overload may disturb the bacteriostatic functions of these proteins and decrease their antimicrobial activity, resulting in increased incidence of infections. Subclinical atherosclerosis in children with major beta-thalassemia begins early in life and these children are at risk of developing premature atherosclerosis.

**Phlebotomy versus Al-hijamah for treating iron overload**

After successful marrow transplantation, iron overload is still an important cause of morbidity in thalassemia. Phlebotomy is a safe, efficient, and widely accepted method where mobilization of iron from overloaded tissues occurs, causing a decrease in iron overload in ex-thalassemic patients. Phlebotomy was reported to decrease serum ferritin and liver iron concentration maximally and significantly. It also improved the histological grading for chronic hepatitis in ex-thalassemic patients. Phlebotomy (versus iron chelation therapy) is still practiced as the therapeutic procedure aiming at getting rid of a portion of blood to excrete an abnormally increased offending blood component. Phlebotomy is the initial treatment of choice in hemochromatosis, while iron chelation therapy is the treatment of choice for transfusional siderosis encountered in thalassemia. A phlebotomy program was reported to be beneficial for two-thirds of patients who underwent allogeneic hematopoietic stem cell transplantation for treating thalassemia. Thalassemia patients (who received bone marrow transplant) with subclinical left ventricular diastolic dysfunction and impaired left ventricular contractility may reverse these processes with an effective regimen of iron reduction using phlebotomy. Interestingly, cure of thalassemia in some hepatic cirrhosis patients with allogeneic bone marrow transplantation resulted in reversibility of liver cirrhosis after iron removal treatment where serum aminotransferase levels decreased, histologic inflammatory activity decreased, and histological biopsies showed regression of incomplete or definite cirrhosis. Phlebotomy may treat iron overload but it may cause significant blood loss, causing aggravation of the anemic status in patients having thalassemia or sickle cell anemia. Blood shed out during phlebotomy may benefit patients by decreasing the level of serum ferritin, iron, antibodies, and hemolyzed cells, but the necessity for a second blood transfusion consequently increases. Elevated erythrocyte ferritin during phlebotomy therapy in iron overload depends on the body iron stores that can be regulated by the activity of erythropoiesis.

However, phlebotomy was not reported to treat the above-mentioned causes of hemolysis in thalassemic patients. As for Al-hijamah, it may treat some causes of hemolysis; eg, through excretion of immunoglobulin antibodies and noxious substances. Al-hijamah-induced percutaneous excretion of iron and ferritin may improve the therapeutic outcomes in iron overload patients (Table 2). Al-hijamah may induce pharmacological potentiation (decreasing the needed dose and frequency of iron chelation drug administration and causing decreased drug-induced side effects). Al-hijamah may enhance the natural immunity and enhance the removal of cellular fragments of hemolyzed cells. Reported WCT-induced blood clearance of antibodies may treat some causes of hemolysis and benefit iron overload patients. Al-hijamah may decrease the needed frequency (and amount) of blood transfusions. Al-hijamah-induced reduction in blood pressure and blood clearance of triglycerides, LDL cholesterol, and total cholesterol may prevent and treat disease conditions associated with thalassemia, such as premature atherosclerosis, and improve the general status in hepatitis patients.

Interestingly, Al-hijamah efficiently treated other diseases, including rheumatoid arthritis, headache, hypertension, and bronchial asthma (Tables 3 and 4). Reported therapeutic benefits of production of endogenous nitric oxide (antioxidant, antimicrobial, and vasodilator) may benefit thalassemia patients. Serum iron, serum ferritin, necessity, and frequency of repeated blood transfusions should be compared before and after Al-hijamah. Al-hijamah was reported to improve the oxygen...
Saturation and respiratory functions better than phlebotomy in cigarette-smoking patients having chronic obstructive pulmonary disease.\textsuperscript{27}

Interestingly, Al-hijamah did not induce anemia or significant blood loss, as evidenced by the normal hematological indices after practicing Al-hijamah.\textsuperscript{27} Al-hijamah (suction, scarification, and suction method) was reported to significantly decrease the level of serum C-reactive protein better than phlebotomy in patients having urticaria and angioedema;\textsuperscript{69} Al-hijamah decreased serum CRP from 11.95 to 1.1 mg/L, while phlebotomy decreased it to a much lower extent.\textsuperscript{69}

### Table 2 Al-hijamah versus phlebotomy for treating iron overload

<table>
<thead>
<tr>
<th>Nature</th>
<th>Phlebotomy</th>
<th>Al-hijamah</th>
</tr>
</thead>
<tbody>
<tr>
<td>Route of excretion</td>
<td>Excretory therapeutic procedure</td>
<td>Percutaneous excretion</td>
</tr>
<tr>
<td>Correcting causes of hemolysis</td>
<td>Venesection</td>
<td>Less powerful</td>
</tr>
<tr>
<td>Medical principles</td>
<td>Removal of a significant portion of whole blood in blood transfusion bags to decrease the concentration of an offending component</td>
<td>Removal of a significant portion of whole blood containing a portion of noxious substances</td>
</tr>
<tr>
<td>Indications</td>
<td>Therapeutic indications</td>
<td>Al-hijamah</td>
</tr>
<tr>
<td></td>
<td>To treat polycythemia and iron overload conditions; eg, thalassemia</td>
<td>Excretory therapeutic procedure</td>
</tr>
<tr>
<td>Side effects</td>
<td>Excessive removal of blood may cause anemia</td>
<td>Repeating the procedure</td>
</tr>
<tr>
<td></td>
<td>Phlebotomy may aggravate anemic state in thalassemia patients</td>
<td>Repeatable according to indication</td>
</tr>
<tr>
<td>Repeating the procedure</td>
<td>Repeatable according to indication</td>
<td>Repeatable according to indication every 2 weeks to 1 year</td>
</tr>
<tr>
<td>Nature of excretion</td>
<td>Whole blood</td>
<td>Whole blood</td>
</tr>
<tr>
<td>Degree of loss of blood cells</td>
<td>Whole blood loss</td>
<td>Whole blood loss</td>
</tr>
<tr>
<td>Steps</td>
<td>Bleeding in a blood collection bag to remove about 500 cc blood</td>
<td>Bleeding in a blood collection bag to remove about 500 cc blood</td>
</tr>
<tr>
<td>Place for receiving treatment</td>
<td>Hematology department in hospitals</td>
<td>Outpatient clinic or home</td>
</tr>
<tr>
<td>Treatment of associated disease conditions</td>
<td>No report</td>
<td>Reported to treat viral hepatitis, hypertension, hyperlipidemia, pain conditions, and others\textsuperscript{1,28}</td>
</tr>
<tr>
<td>Other therapeutic benefits</td>
<td>None</td>
<td>Reported to treat viral hepatitis, hypertension, hyperlipidemia, pain conditions, and others\textsuperscript{1,28}</td>
</tr>
<tr>
<td>Pharmacological potentiation effect</td>
<td>No report</td>
<td>More selective (loss of lesser amount of the red blood cell mass)</td>
</tr>
<tr>
<td>Duration of the procedure</td>
<td>Few minutes</td>
<td>Skin suction, scarification and second suction (triple S technique)</td>
</tr>
<tr>
<td>Clearance of skin iron</td>
<td>None</td>
<td>Skin suction, scarification and second suction (triple S technique)</td>
</tr>
<tr>
<td>Other names</td>
<td>Fasd (in Arabic), bloodletting</td>
<td>Reported\textsuperscript{1}</td>
</tr>
<tr>
<td>Separation of noxious substances from blood</td>
<td>Not done</td>
<td>0.5–1 hour</td>
</tr>
<tr>
<td></td>
<td>There is only excretion of a significant portion of blood containing a portion of noxious substances</td>
<td>Yes</td>
</tr>
<tr>
<td>Extent of improvement</td>
<td>Depends on the amount of letted blood (containing noxious substances) versus exaggerating the anemic state</td>
<td>Depends on the amount of excreted noxious substances with proper pressure-dependent filtration and capillary blood clearance (skill of practice). No exaggeration of the anemic condition</td>
</tr>
</tbody>
</table>

**Abbreviation:** WCT, wet cupping therapy.

#### Reported therapeutic benefits of Al-hijamah during treatment of iron overload

There are a lot of reported hematological health benefits that can be gained when treating patients having different diseases with Al-hijamah. However, the most attractive reported side effect of repeated traditional WCT is iron deficiency anemia, which can be utilized for treating iron overload conditions. Practicing Al-hijamah was not reported to cause massive blood loss or anemia. Instead, Al-hijamah was reported to reduce serum ferritin significantly.\textsuperscript{1} Although there are no published reported studies about the use of Al-hijamah for treating...
Table 3 Al-hijamah versus iron chelation therapy for treating iron overload

<table>
<thead>
<tr>
<th>Examples and nature of treatment</th>
<th>Iron chelation therapy</th>
<th>Al-hijamah (triple S technique)</th>
</tr>
</thead>
</table>
|                                  | • Deferoxamine, deferasirox and deferiprone  
• Pharmacological treatments  
• Better to be combined with Al-hijamah for treating iron overload  | • Al-hijamah is a simple percutaneous excretory procedure that is distinct from traditional WCT  
• Better to be combined with iron chelation therapy  |

| Route of administration (or method of practice)\(^{95-98}\) | For deferoxamine:  
Subcutaneous or intravenous as continuous infusion 5–7 days weekly; not orally available; 20–60 mg/kg/day averaged over a week if not given daily  
For deferasirox:  
Oral, as a suspension, once daily (or in some circumstances divided twice daily); 20–40 mg/kg/day, highly individualized and is dependent on the transfusion rate  
For deferiprone:  
Oral as tablets, generally in three divided doses; 75 mg/kg/day  | Percutaneous  |

| Mechanism of action\(^{95-98}\) | For deferoxamine:  
Binds free iron in the blood to enhance its urinary excretion  
Removes excess tissue iron; eg, the liver  
Affects expression and release of inflammatory mediators by specific cell types\(^{60,61}\)  
For deferasirox:  
Selective for iron (as Fe\(^{3+}\))  
Binds iron with high affinity in a 2:1 ratio  
For deferiprone:  
Has an affinity for ferric ion (iron III). Deferiprone binds with ferric ions to form neutral complexes that are stable over a wide range of pH values  | Pressure-dependent, size-dependent nonspecific filtration of blood circulation through the fenestrated dermal capillaries causing nonspecific blood clearance;\(^{1,18}\) eg, excretion of iron and ferritin  |

<table>
<thead>
<tr>
<th>Nature of iron excretion</th>
<th>Pharmacological iron excretion</th>
<th>Physiological pressure-dependent mechanism</th>
</tr>
</thead>
</table>

<table>
<thead>
<tr>
<th>Methodology</th>
<th>Indirect iron excretion</th>
<th>Direct iron excretion</th>
</tr>
</thead>
</table>

| Clearance of blood and interstitial fluids | Reported to clear blood of excess iron and ferritin  | Reported to clear both blood and interstitial fluids from excess pathological substances; eg, autoantibodies and ferritin in a nonspecific manner\(^{1,11}\)  |

| Indications | Iron overload conditions  | A long list of disease conditions that include pain conditions (eg, back pain), autoimmune diseases (eg, rheumatoid arthritis), neurological conditions (eg, headache), infections (eg, cellulitis), and others\(^{1-3,13,63}\)  |

<table>
<thead>
<tr>
<th>Route of iron excretion</th>
<th>Urine and stool</th>
<th>Percutaneous</th>
</tr>
</thead>
</table>

| Frequency of administration | Daily  | Every 1–3 months  |

<table>
<thead>
<tr>
<th>Tolerability</th>
<th>Tolerable in most cases except when allergy or severe side effects develop</th>
<th>Tolerable</th>
</tr>
</thead>
</table>

| Duration per a single treatment | For deferoxamine, Infusion takes about 8 hours per session  
For deferasirox and deferiprone, treatment is oral  | 0.5–1 hour  |
|-----------------------------|---------------------------------------------------------------|-------------|

| Plasma half-life\(^{95-98}\) | For deferoxamine: Short (~20–30 minutes)  
For deferasirox: Long (11–16 hours)  
For deferiprone: Intermediate (~2–3 hours)  | No half-life as Al-hijamah is a minor surgical excretory procedure  |

| Other therapeutic benefits | None  | Pharmacological potentiation, immunological potentiation, analgesic effect, improvement of microcirculation, and others  |

| Therapeutic values of combining Al-hijamah with iron chelation therapy | Al-hijamah-induced excretion of iron and ferritin may:  
• Improve the therapeutic effects of iron chelators  
• Decrease the needed drug doses  
• Decrease the frequency of drug administration  
• Decrease the duration of scheduled treatment  
• Decrease the drug-induced side effects  | Combining iron chelators with Al-hijamah may:  
• Potentiate the state of negative iron balance (Fe output \(>\) Fe input)  
• Mobilize more iron from tissues to blood  
• Enhance iron excretion  
• Protect vital organs from tissue damage  |

| Treatment of other diseases or associated disease | None  | Yes. WCT and Al-hijamah were reported to treat many disease conditions that are different in etiology and pathogeneses  |

Abbreviation: WCT, wet cupping therapy.
thalassemia or conditions of iron overload, the safety of practicing Al-hijamah and the reported health benefits associated with it may be an attractive topic for hematologists and researchers to explore. The therapeutic benefits of Al-hijamah for treating different disease conditions have been previously reported. Many of those health and therapeutic benefits are helpful when treating patients having iron overload with hemolysis; for example, thalassemia. Al-hijamah-induced immunological enhancement (eg, physiological leukocytosis and increased count of NK cells) is beneficial for improving the poor immunity status in thalassemia patients. Blood and interstitial fluid clearance from excess ferritin and iron is the single most important therapeutic benefit of Al-hijamah when treating thalassemia patients. Being a percutaneous excretory procedure, skin (a site for iron deposition) clearance of excessive iron can be gained through Al-hijamah. Moreover, nonspecific blood clearance gained during Al-hijamah, where patients’ serum can be cleared significantly of LDL cholesterol, total cholesterol, and triglycerides, may protect thalassemia patients against the development of premature atherosclerosis. Pharmacological potentiation reported when combining Al-hijamah with different drug treatments

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**Table 4 Safety issue of iron chelation therapy versus Al-hijamah**

<table>
<thead>
<tr>
<th>Iron chelation therapy</th>
<th>Al-hijamah (SSS technique)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Safety of medical practice</td>
<td>Administration of deferoxamine, deferasirox, and deferiprone is usually safe but has certain limitations, side effects, contraindications, and precautions</td>
</tr>
<tr>
<td>Special precautions</td>
<td>For deferasirox: Acute renal failure and cytopenias; eg, agranulocytosis, neutropenia, and thrombocytopenia For deferiprone: Hepatic and renal impairment Neutropenia</td>
</tr>
<tr>
<td>Contraindications</td>
<td>For deferiprone: Agranulocytosis, pregnancy, and lactation</td>
</tr>
<tr>
<td>Known or theoretical disadvantages</td>
<td>For deferoxamine: • Frequent parenteral administration may not be tolerated • Local skin reactions</td>
</tr>
<tr>
<td></td>
<td>For deferiprone: • Gastrointestinal or joint symptoms • Agranulocytosis</td>
</tr>
<tr>
<td></td>
<td>For deferiprone: • Neutropenia require weekly CBC monitoring</td>
</tr>
<tr>
<td></td>
<td>For deferasirox: • Gastrointestinal symptoms</td>
</tr>
<tr>
<td>Common side effects</td>
<td>For deferoxamine: Local skin reactions; sensorineural hearing loss, and bone problems For deferasirox: Rash gastrointestinal upset, diarrhea, mild abnormalities in creatinine clearance and proteinuria For deferiprone: Gastrointestinal distress, joint pain, and erosive arthritis and neutropenia</td>
</tr>
<tr>
<td>Severe and/or dangerous side effects</td>
<td>For deferoxamine: Siderophore for some bacteria (eg, listeria) Retinopathy and acute pulmonary disease For deferasirox: Peptic ulcers, liver dysfunction, renal dysfunction (including failure), cytopenias For deferiprone: Agranulocytosis and hepatic fibrosis</td>
</tr>
<tr>
<td>Drug interactions</td>
<td>Avoid using deferiprone with aluminum-containing antacids, as it can chelates trivalent metal ions</td>
</tr>
</tbody>
</table>

**Abbreviations:** CBC, complete blood count; SSS, triple S technique; WCT, wet cupping therapy.
may enhance the therapeutic effects and decrease the side effects of concomitantly administered drugs by iron overload patients, such as iron chelation therapy. Importantly, Al-hijamah was not reported to cause massive bleeding (red blood cell loss)\textsuperscript{36,37} or aggravate anemic status. Al-hijamah induces possible nonspecific blood clearance of liberated hemoglobin and fragments of hemolyzed cells (having small particle sizes that can be filtered through capillary pores).

**Al-hijamah combined with oral honey for treating hyperferremia and its complications**

Consumption of oral honey was reported to provide numerous beneficial effects that may expand the therapeutic effects of Al-hijamah when combined with Al-hijamah. Oral honey may benefit thalassemia and iron overload patients, especially thalassemia cases complicated with diabetes mellitus, poor immunity, and oxidative stress-induced effects; oral honey improved body weight and serum lipid profile of diabetic patients\textsuperscript{38} to the extent that it was recommended to be used as a sugar substitute in patients with type 1 diabetes mellitus\textsuperscript{71} or impaired glucose tolerance.\textsuperscript{72} Patients having impaired glucose tolerance exhibited significant lower plasma glucose concentrations after consumption of oral honey where the plasma glucose levels peaked at 30–60 minutes in response to oral honey and showed a rapid decline as compared to patients consuming glucose, which indicated a better control of the glycemic index with oral honey administration.\textsuperscript{72} Oral honey consumption was reported to increase serum antioxidant capacity, vitamin C concentration, beta-carotene, and glutathione reductase. Interestingly, honey reduced plasma ferritin by 11%.\textsuperscript{73} Moreover, oral honey was reported to enhance the natural immunological response through increasing the monocytes and lymphocytes percentage, decreasing serum immunoglobulin E\textsuperscript{73} and stimulating antibody production during primary and secondary immune responses against thymus-dependent and thymus-independent antigens.\textsuperscript{74}

Interestingly, oral honey ingestion was reported to decrease prostaglandins levels, elevate nitric oxide production, and improve the hematological and biochemical tests in a patient with a long history of AIDS.\textsuperscript{75} It might be concluded that combining Al-hijamah with oral honey may improve the therapeutic outcomes in thalassemia and iron overload patients. Interestingly, in prophyptic medicine, both Al-hijamah and oral honey are curative remedies according to the prophetic teaching: “Cure is in three: in sharat mihjam, gulp of honey (oral honey), and cauterization. I do not recommend my nation to cauterize.”\textsuperscript{76,77}

**Conclusion**

In conclusion, Al-hijamah is a beneficial, safe, economic, time-saving (about 30 minutes), and repeatable treatment modality. Al-hijamah may be a good combinatory treatment with iron chelation therapy, and can be a sole treatment when there is a limitation or contraindication to iron chelation therapy. Al-hijamah may be superior to phlebotomy, and it needs to be researched further for educational evaluation in treating iron overload. Oral honey also carries a lot of therapeutic health benefits when treating conditions of iron overload. This may strongly suggest Al-hijamah combined with oral honey as a novel treatment for thalassemia and iron overload conditions may potentiate therapeutic effects of iron chelation therapy.

**Acknowledgments**

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**Disclosure**

The authors declare no conflict of interest in this work.

**References**


Al-hijamah for thalassemia, iron overload, and hyperferremia