Clinical study of Ghritkumari (Aloe-vera) and Kakamachi (Solanum nigrum) in Kamala roga (Jaundice)

Authors
Dinbandhu Kumar Kanavjiya¹, Dr Vijay Shankar Dubey², Dr Amrendra Kumar Singh³
¹MD (Ay), AMO, GGSH., Patna, 7004898087
²Prof. Dravyaguna Dept., Govt. Ayurveda college Patna
³Asso. Prof., Rog evam Vikriti Dept., Govt. Ayurveda College Patna

According to Indian philosophical traditions it is believed that Ayurveda has a divine origin that is, it has been created by Brahma the creator of universe and has been transmitted to various saints and scholars to remove the sufferings and miseries of all living subjects throughout the world. The present study entitled “Gunakarmatmak study of Ghritkumari (Aloe vera) and Kakamachi (Solanum nigrum) with special reference to Kamala (Jaundice)” has been undertaken to launch a literary and clinical study of Kamala (Jaundice) along with therapeutic evaluation of the efficacy of Ghritkumari leaf and Kakamachi Panchang churna in patients of Kamala. Ayurveda is really well known for its simple, promotive, preventive, curative and holistic approach. Ghritkumari and Kakamachi which are very useful Ayurvedic drug, one of the most easily available, popular, widely utilized and described by all ancient Acharyas like Charaka, Sushruta, Vagbhata and all Nighantu Kalas have been selected for the present research work. Here, the drug Ghritkumari and Kakamachi has been comprehensively evaluated by all aspects regarding it.

Drug Review Historical Background
Aloe vera: In the foremost ever written document of knowledge, we cannot trace any reference regarding the drug Kumari. In Bhela Samhita, Kumari is mentioned as an ingredient of a compound in the treatment of Vata Vyadhi. There is no any reference found in Brihatrayi. In Sharangadhara Samhita, its use is described in Plilha Roga. All Nighantu Kalas have mentioned Kumari in different Vargas. In 1932 it was included in British pharmacopoeia. But in Ayurvedic amenities there is no any direct reference of Kumari regarding as Yuvanapidika.
Kakamachi: It is not mentioned in the Vedic literature. In Charaka Samhita references regarding Kakamachi is present in Sutrasthana, Nidanasthana, Vimanasthana and also Chikitsasthana. Kakamachi is highlighted for its Kustaghna property, in Charaka sutrasthana 3/16.

Again in Atryebhadrakapyeeya adhyaya – 84 sloka, Charaka mentions Kakamachi in the context of Virudha aharas by saying that, Kakamachi used after a long time, after cooking is Virudha. Aloe vera- About 14 synonyms have been traced regarding Ghritkumari in various
samhita granths and Nighantus. The synonyms of Ghritkumari to morphological characters in correlation with that of botanical description of the plant Aloe vera. The synonyms like Ghritkumari, Kumari, Grhakanya, Visala, Mndala, Bahuptra etc. indicate the morphological characters. Kakamachi- As well as synonyms of Kakamachi are Kakamata, Gudaphala, Kakini, Vara, Sundari, and Gucchaphala etc. **Rasapancaka**: ALOE VERA - According to Ayurvedic pharmacology, the drug is based on Rasa Pancaka viz. Rasa, Guna, Virya, Vipaka and Prabhava. In Brhatrayi Acaryas stated that some of the drug act is accordance with their taste, some with properties, some with potency, some with post digestive effect and some with specific action. (Ch. Su. 26/17, Su. Su.40/17, AS.Su. 17/30). Ghritkumari has been established Guru, Snigdha, Pichila in Guna; Tikta and Madhura in Rasa; Katu in Vipaka, sita in Virya. According to Doshaghnata it is Tridosasamakka. **KAKAMACHI** - Kakamachi has been established Laghu, Snigdha in Guna; Tikta in Rasa; Katu in Vipaka, Anusna in Virya. According to Doshaghnata it is Tridosaghna.

**Therapeutic Actions**: Aloe vera: Kumari has been important and potent drug in ancient medicine and therapeutics finding its uses in several diseases as mentioned in different classical texts dealing with clinical management. Kamala (jaundice) by Bhavaprakasha, Pandu (anemia) Due to pharmacological actions on liver and spleen. Juice of Kumari is given in Yakrta and Plhav rvidhi and roga. Due to poten cholagogue (Pittanirharana, Pittarecana). The juice of Kumari mixed with Haridra (turmeric) powder is given in spleen enlargement or Plhav rviddi. by Sharangadhar Samhita. Rajaoabroda (amenorrhoea) Kumarikavati and Rajah pravartani vati are prescribed in menstrual problems as emmenagogue drugs by Bhaishajya Ratnavali. Extract of Kumari leaves juice (Kanyasara) is given. Varna (wound) and abscess Kumari mixed with tila and sour gruel or alone is applied to ripens the abscess. Kumari leaves or juice is recommended to apply over abscess and wounds in ama and pacyamana stages for attaining pakva condition of vidradhi and vrana. by Vaidya Manorama. Leaves pulp of Kumari are prescribed to externally apply over or by covering up with the steamed leaves devoid of pulp, in all stages of abscess as indicated in medical texts. by Vidradhi cikitsa context Kakamachi:- It is told that Kakamachi is Tridoshashamaka, Snigdha, Ushna, Tikta, Rasayani, Swary, Vrushya etc. and acts against Yaktra viridhi, Shotha, Kusta, Arsha, Jvara, Meh, Hikka, Chadri, and Hrudroga.

**Therapeutic Indication**: Ghritkumari is mostly indicated in the diseases like Agnidagdha, Raktavarakshotha, Dourbalya, Krimiogatant-ukrini, Yaktrplihavikara vriddhi, PanduKamala Modern authors indicate it in, Jaundice, Anticancer activity, cough drops, Analgesic: Antiinflammatory activity etc. CULTIVATION: - Soil: Naturally occurs in driest and poorest soils and can be grown in variety of soil. But the most ideal soil for its sandy loam that is slightly alkaline with a PH up to 8.5. Theroot system of this plant is shallow and does not penetrate deep into the soil. However, water logged soil is totally unsuitable.

**Pharmacognostical Study**: Pharmacognostical study of plant drugs plays a major role in the standardisation and quality control of the plant drugs. The study of macroscopic and microscopic characters was carried out. Morphology: Aloe vera: A perennial and coarse looking plant is with a short, thick, cylindrical, simple, woody stem, 1-2 feet high, sending out at the base numerous stolons. Kakamachi: Herbs or shrubs, erect, trailing or scandent, rarely small trees. Leaves are alternate often in unequal pairs rarely clustered never truly opposite, entire, lobed or pinnate. Flowers regular hermaphrodite, terminal or lateral axillary or extraaxillary cymes or solitary or clustered pedicels.

**Phytochemical Study**: Phytochemical studies play a very important role in the standardization of any single or compound drug. The loss on drying, Ash Value, Water soluble extract and
Disintegration time of Ghiritkumari and Kakamachi were of the standard level with comparison to Ayurvedic pharmacopoeia of India. Another data regarding UV spectra test and T.L.C. study of the sample are also compared with the same which is deeply discussed in the section of phytochemical study. As the phytochemical analysis of the Ghiritkumari and Kakamachi could not be compared due to lack of comparative data in the absence of any previous study on it. But obtained data from the various test like, Refractive index, Specific gravity, Acid value, Iodine value and Saponification value of Ghiritkumari and Kakamachi would be helpful to analyzed and to standardize the drug for the next study. Different types of chemical constituents like Anthraquinones, Lignin, Saponins, Fatty acids, Salicylic acid, and Amino acids of Aloe vera. TLC results indicate Musabbar (Alua) Rf value of spots a visualized in UV 366 nm at Rf. 0.10, 0.30, 0.40, 0.60 & 0.75 (5 spots). Rf value of spots visualized in UV 254 nm at Rf. 0.24, 0.30, 0.40, 0.50, 0.55 & 0.75(6) Rf value of spots visualized after spray of sulphuric acid reagent and heated 110°C for 5 minutes at Rf. 0.20, 0.30, 0.55, 0.75 (4 spots). Leaves contain quercetin glycosides. Immature fruits contain gluco alkaloids. Leaves and fruits contain solasodine type compounds. Fruits contain steroidal glycosides, glycoalkaloids, α – solamargine & α – solasonine. Seeds contain fatty oil of Kakamachi. TLC results indicate Kakamachi Rf value of spots visualized in Iodine at Rf 0.10, 0.30, 0.40, 0.60 & 0.75(6 spots). Rf value of spots visualized in UV 254 nm at Rf. 0.24, 0.30, 0.40, 0.50, 0.55 & 0.75 (6 spots) Rf value of spots Visualized after spray of sulphuric acid reagent and heated 110°C for 5 minutes at Rf 0.10, 0.25, 0.30, 0.35, 0.40, 0.60, 0.65, 0.75 & 0.85 (9 spots).

**Disease review:** 1. “Kayam malayte iti Kamala” It means the whole body becomes dirty, because of the accumulation of mala i.e. mala ranjaka pitta. 2. Kama + La = Kamala. Kamam Lati Hanti Iti Kamala. It means there is aversion for all desires in the patient of Kamala which may because of his physical and mental disability that is seen in this disease. “Kutsita mala yasmin Roga sah Kamala Roga.” Regularly produced mala and its excretion from the body is essential phenomena maintain to the health. As this function has been enunciated by Charaka and Susruta. “Dosha Dhatu mala moolam hi shariram.” The word Jaundice has been derived from the French word “Jaundice” which means yellow discolouration of the body organs. In Ayurveda similar nomenclature has been given for so many diseases on the basis of symptoms like Pandu, Prameha, and Atisara etc. But in the case of this disease they have preferred to name it as Kamala, which is based on pathogenesis. It is to indicate that in this disease, the whole body is highly vilated by male ranjaka pitta (Bilirubin) which is the by product of rakta dhatu. TYPES OF KAMALA: Many classification of kamala have been suggested by various authors. They have classified kamala in one way or the other way, as described in different Ayurvedic classics. Charaka has mentioned two types of kamala (Ch.Su.19/4). (1) Koshtashritra, (2) Shakhashritra Kamala. But only koshtashritra Kamala is a misnomer. In Kamala pitta roopa mala has to spread all over the body may be skin, eyes, mouth etc. So the word koshtashritra has been introduced in place of koshtashritra kamala and the same has been referred in Chikitsa sthana and other places. Of course shakhashraya Kamala is observed in rudhapatha kamala where due to obstruction of the biliary passages there is no pouring of pitta in Kamala so it means it gives rise to shakhashritra Kamala. The word shakha here refers to skin, rakta, mamsa etc. Thus appropriately the two types of Kamala should be referred as follows: (1) Koshtashakhshritra Kamala (2) Shakhshritra Kamala.

**Clinical Study**- The clinical study was carried out on 30 patients of Kamala (Jaundice), which were divided into 3 groups viz. Group I Application of Ghiritkumari. Group II Application of Kakamachi. Group III Combined group. The patients were selected randomly irrespective of age, sex, marital status, Education etc. The observation and results
obtained are discussed here. For the assessment of results of the therapy, the patients were examined subjectively. Signs and symptoms were assessed by adopting suitable scoring methods as mentioned in materials and methods. The results obtained were statically analyzed and mean change in %, S.D, ‘t’value, p value were calculated for each criteria of assessment in all group.

Selection of patients: For the present study, patients fulfilling the clinical criteria for diagnosis of Kamala (Jaundice) were selected irrespective of their age, sex, religion etc, in random from O.P.D. and I.P.D. section of Govt. Ayurvedic College and Hospital, Kadam kuan, Patna.

Preparation of Drug: Drugs were prepared in Dravyaguna Pharmacy of Govt. Ayurvedic College and Hospital, Kadam kuan, Patna.

Incidence: 1. Age: Group I: Majority of the patients were of age group 21-40 yrs – 5 (50%) followed by Gr. 41-60 yrs 4 (40%), 12-20 yrs 1(10%). Group II: Majority of the patients were of age group Group II: p 21-40 yrs – 5 (50%) followed by Gr. 12-20 yrs 3 (30%), Gr. 41-60 yrs – 2(20%) Group III: Majority of the patients were of age group 21-40 yrs – 7 (70%) followed by Gr. 12-20 yrs 2 (20%), Gr. 41-60 yrs – 1(10%). Overall incidence of Majority of patients were of age Group 21-40yrs 17(56.66%) followed by age Gr.41-60 yrs (23.33%) and least in age group 12-20 yrs(20%). In present study maximum 17 patients out of 30 were between 21-40 yrs. (2) Sex: Group I: Group I: Majority of patients were male i.e. 8 (80%) and female patients were 2 (20%) Group II: Majority of patients were male i.e. 7 (70 Group II: %) and female patients were 3(30%) Group III Majority of patients were male i.e. 7 (70%) and female patients were 3(30%) (3) Marital status: Group I: Majority of patients were unmarried i.e. 9 Group I: (90%) and married patients were 1 (10%). Group II: Majority of patients were married i.e. 5 Group II: (50%) and unmarried patients were 5 (50%). Group III Majority of patients were married i.e. 5 (50%) and unmarried patients were 5(50%) However any consolidated conclusion may not be drawn from the findings of this data, even then 90% patients were unmarried and due this they were conscious about their outfit and reported in more number at the clinic. 4. Educational profile Group I Majority of patients had matric 7 (70%), Group I patients with Graduate 2 (20%), matric 1(10%), primary and illiterate 0 (0%). Group II Majority of patients had Inters 3 (30%), patients with Graduate 2 (20%), primary 2(20%) and illiterate 2(20%). Primary 1(10%) Group III Majority of patients had Graduate 4 (40%), Inter3 (30%), patients with, matric 2 (20%) and illiterate1 (10%).Primary 1 (0%) Though educational qualification does not have any direct effect on the occurrence of Kamala, it may be concluded that education creates awareness in the individuals to seek proper cure for their disorders and consult the physician specialized in the disorders by which they are suffering from. Matric the predominant age group was study period thus higher percentage (40%) belongs to educational group. 5. Occupation Group I Majority of patients were businessman 4 (4 Group I 0%), farmer 2 (20%), House wife 2(20%), serviceman 1(10%) and student 1(10%). Group II – Majority of patients were student 4 (40%), serviceman 2 (20%), House wife 2(20%), farmer 1(10%) and businessman 1(10%) Group III Majority of patients were student 5 (50%), House wife 2 (20%), businessman 2 (20%), serviceman 1 (20%), and farmer 0(0%) Almost all the types of occupational groups had the patients of Kamala where the highest number was recorded in the patients who were student farmer or Housewife. The over increasing tension of studies followed by avertiing sleep till late night for reading and due to constant worries, irregular food habits, over indulgence in sleep, addictions like tea, tobacco etc. aggravating the Vata and Pitta Doshas create Kamala in a long run. 6. Food Habitat:: Group I Majority of patients were nonvegetarian 8 Group I (80%), and vegetarian 2(20%). Group II Majority of patients were nonvegetarian 8 Group II (80%), and vegetarian 2(20%). Group III Majority of patients were nonvegetarian 8 (80%), and vegetarian 2 (20%). This may show that Kamala is more
common in non vegetarian. 7. Prakriti:- Group I Majority of patients were vatapitta 8 (8 Group I 0%), vatakapha 1 (10%) and pittakapha 1(10 %).
Group II Majority of patients were vatapitta 7 (70%), vatakapha 2 (20%) and pittakapha 1(10 %). Group II Majority of patients were vatapitta 7 (70%), vatakapha 2(20%) and pittakapha 1(10 %).
A maximum no. of patients belongs to Vata Pitta Prakriti (73.33%) followed by vata Kapha Prakriti (16.66%). Vata and Pitta and are the two main doshas, which play an important role in the occurrence of Kamala (jaundice). It may be concluded from this finding that when such individuals indulge in etiological factors they are more prone to have Kamala. Observations about prakriti are in accordance with textual references.
8. Kostha. Group I Majority of patients were krura 5 (50%) Madhya 3 (30%) and mridu 2(20%).
Group II Majority of patients were krura 7(70%) Madhya 2 (20%) and mridu 1(20%). Group III Majority of patients were krura 7(70%) Madhya 3(30%) and mridu 0(0%) Maximum numbers of patients (63.33%) were found Krura Kostha followed by Madhya Kostha (26.66%). This also indicate the causatively of Vata and Pitta dominated Dwandvaja Prakriti may be responsible for Kamala. 9. S.Billirubin Group I Majority of patients were level of >2 -4 is 4 (40%) followed by level of 1 to 2 is 3 (30%), and level of >4-10 is 3 (30%). Group II Majority of patients were level of 1 to 2 is 7(70%) followed by level of >2-4 is 2 (20%), and level of >4-10 is 1 (10%). Group III Majority of patients were level of >2-4 is 4 (40%) followed by level of 1 to 2 is 3 (30%), and level of >4-10 is 3 (30%). 10. S.G.P.T. Group I Majority of patients were level of >35-100 is 10(100%) followed by level of Up to 35 is 0 (0%), and level of >100 is 0 (0%).Group II Majority of patients were level of >35-100 is 7(70%) followed by level of Up to 35 is 2 (20%), and level of >100 is 1(10%). Group III Majority of patients were level of >35-100 is 8(80%) followed by level of up to 35 is 0(0%), and level of >100 is 0(20%).
Clinical features: In the present clinical study the observation of various cardinal features in patients are as under. Yellowish discoloration eyes and urine. Loss of appetite, Abdominal pain, Nausea/Vomiting, Fever, Muscles cramp, Itching and Weakness.

Effects of Therapy
(A)Effect on symptoms: 1. Yellowish colour of eyes: Group I The mean grade of Yellowish colour of eyes before treatment was 2.3 and it lowered down to 0.5 with SD± 1.032, giving a relief of 78.26 % with ‘t’ value 5.51 (p<0.001) which was statistically highly significant. Group II The mean grade of Yellowish colour of eyes before treatment was 2.4 and it lowered down to 0.6 with SD± 0.632, giving a relief of 75% with ‘t’ value 9 (p<0.001) which was statistically highly significant. Group III The mean grade of Yellowish colour of eyes before treatment was 2.5 and it lowered down to 0.4 with SD± 0.316, giving a relief of 84% with ‘t’ value 21 (p<0.001) which was statistically highly significant.
2. Yellowish colour of urine: Group I The mean grade of Yellowish colour of urine lowered from 2.1 to 0.5 with SD± 0.843 giving a relief of 76.19% with‘t’ value 6 (p<0.001) which was statistically highly significant. Group II The mean grade of Yellowish colour of urine lowered from 2.5 to 0.7with SD± 0.674giving a relief of 72 % with‘t’ value 9 (p<0.001) which was statistically highly significant. Group III The mean grade of Yellowish colour of urine lowered from 2.4 to 0.4 with SD± 0.471 giving a relief of 83.33 % with’t’ value 13.41 (p<0.001) which was statistically highly significant.
3. Loss of appetite: Group I The mean grade of loss of appetite lowered from the pretrial value of 2.5 to 0.4 with SD±0.918, giving a relief of 72% with't’ value 6.19 (p<0.001) which was statistically highly significant. Group II The mean grade of loss of appetite lowered from the pretrial value of 2.3 to 0.7with SD±0.699, giving a relief of 69.56% with’t’ value 7.23 (p<0.001) which was statistically highly significant. Group III The mean grade of loss of appetite lowered from the
pretrial value of 2.6 to 0.2 with SD± 0.516, giving a relief of 92.3% with ‘t’ value 14.69 (p<0.001) which was statistically highly significant.

4. **Abdominal pain**: Group I The mean grade of abdominal pain lowered from 0.8 to 0.2 with SD± 0.843, giving a relief of 75% with ‘t’ value 2.25 (p>0.02) which was statistically not significant. Group II The mean grade of abdominal pain lowered from 1.4 to 0.4 with SD± 0.816, giving a relief of 71.42% with ‘t’ value 3.87 (p<0.01) which was statistically significant. Group III The mean grade of abdominal pain lowered from 1.33 to 0 with SD± 0.51, giving a relief of 100% with ‘t’ value 6.32 (p<0.001) which was statistically highly significant.

5. **Nausea/Vomiting**: Group I The mean grade of nausea/vomiting lowered from 1.1 to 0.2 with SD± 0.276, giving a relief of 81.81% with ‘t’ value 3.25 (p<0.02) which was statistically significant. Group II The mean grade of nausea/vomiting lowered from 1.8 to 0.5 with SD± 0.674, giving a relief of 72.22% with ‘t’ value 6.09 (p<0.001) which was statistically highly significant. Group III The mean grade of nausea/vomiting lowered from 1.8 to 0.2 with SD± 0.547, giving a relief of 88.88% with ‘t’ value 6.53 (p<0.001) which was statistically highly significant.

6. **Fever**: Group I The mean grade of Fever lowered from 0.8 to 0 with SD± 1.032, giving a relief of 100% with ‘t’ value 2.44 (p>0.02) which was statistically not significant. Group II The mean grade of Fever lowered from 1.1 to 0.4 with SD± 0.674, giving a relief of 63.63% with ‘t’ value 3.27 (p<0.01) which was statistically significant. Group III The mean grade of Fever lowered from 1.2 to 0.3 with SD± 0.5, giving a relief of 100% with ‘t’ value 5 (p<0.001) which was statistically highly significant.

7. **Muscles cramp**: Group I The mean grade of muscles cramp from 0.8 to 0 with SD± 1.032, giving a relief of 100% with ‘t’ value 2.44 (p>0.02) which was statistically not significant. Group II The mean grade of muscles cramp from 1.1 to 0.3 with SD± 0.788, giving a relief of 72.72% with ‘t’ value 3.20 (p<0.02) which was statistically significant. Group III The mean grade of muscles cramp from 1.5 to 0 with SD± 0.707, giving a relief of 100% with ‘t’ value 3 (p<0.01) which was statistically significant.

8. **Itching**: Group I The mean grade of itching lowered from 0.2 to 0 with SD± 0.632, giving an improvement of 50% with ‘t’ value 1 (p>0.10) which was statistically not significant. Group II The mean grade of itching lowered from 1.4 to 0.7 with SD± 0.674, giving an improvement of 50% with ‘t’ value 3.27 (p<0.01) which was statistically significant. Group III The mean grade of itching lowered from 1 to 0.5 with SD± 0.577, giving an improvement of 50% with ‘t’ value 1.7 (p>0.1) which was statistically not significant.

9. **Weakness**: Group I The mean grade of weakness lowered from 2.6 to 1 with SD±0.51, giving an improvement of 61.53% with ‘t’ value 9.79 (p<0.001) which was statistically highly significant. Group II The mean grade of weakness lowered from 1.2 to 0.6 with SD±0.699, giving an improvement of 50% with ‘t’ value 2.71 (p>0.02) which was statistically not significant. Group III The mean grade of weakness lowered from 2.4 to 0.3 with SD±0.567, giving an improvement of 87% with ‘t’ value 11.69 (p<0.001) which was statistically highly significant.

**Effect of Laboratory Parameters**

For the diagnosis and assessment of results, the main laboratory criteria were serum bilirubin (B.total) estimation. Group I The mean grade of billirubin total lowered from 2.97 to 1.40 with SD± 0.955, giving a relief of 52.93% with ‘t’ value 5.21 (p<0.001) which was statistically significant. Group II The mean grade of billirubin total lowered from 1.2 to 0.6 with SD±0.699, giving an improvement of 50% with ‘t’ value 2.71 (p>0.02) which was statistically not significant. Group III The mean grade of billirubin total lowered from 1.96 to 1.66 with SD± 0.38, giving a relief of 15.29% with ‘t’ value 2.49 (p>0.02) which was statistically not significant. Group III The mean grade of billirubin total lowered from 3.869 to 1.131 with SD± 2.01, giving a relief of 70.76% with ‘t’ value 4.29 (p<0.01) which was statistically significant.
Sides Effects of Therapy
In the present clinical study, two of the registered patients reported adverse effect of the drug group III (Ghiritkumari and Kakamachi) during the therapy. Two patients reported that after taking these groups III drugs them suffering from heartburn. Haematological investigations carried out before and after the treatment showed no major change. The above findings suggest that there is no systemic ill effect of the drugs Ghiritkumari and Kakamachi. But the Drugs needs to be further evaluated for ill effects on large samples so that the drug can be prescribed for longer duration with complete safety.

Probable Mode of Action
The present clinical study reveals that Ghritkumari and Kakamachi used are effective in the treatment of Kamala. The results were more appreciable in group III, in both the clinical and laboratory criteria. Statistically, relief in Yellowish color of eyes and urine, loss of appetite, weakness and in reduction in serum bilirubin levels was highly significant. The trial drugs were Ghritkumari and Kakamachi have been used in management of Kamala (Jaundice) in Ayurvedic texts and Nighantus. Experimental studies have shown them to be improves appetite and taste, Moothrala, Yakrituttejaka, moothrala properties. Ghritkumari is Vatahara, Kaphahara and Tridoshhara. Kakamachi is Tridosha Shamak especially kapha shamak. Therefore combination of both the drugs are more significant than individual. Once the doshik homeostasis has been achieved, the signs and symptoms of Kamala are relieved automatically because the disease and its different manifestations are all produced by doshas. Ghritkumari – Ghritkumari is Tikta, Madhur in rasa, Shita Virya. Guru, Snigdha and Pichcchila Guna.It is said of Vatahara, Kaphahara and Tridoshhara shamaka, Bhedana, Krimighana, Kushtthaghan, Balya, Rasyana, Virecan, Dipan Pacana and Rechana. It is mainly indicated in Pandu, Kamala, Yakritiplha vikaravrddhi, Dourbalaya, Agnidagdha roga etc. Ghritkumari is Tikta, Madhur in Rasa. Katu in Vipak and Guru, Snigdha, Pichcchila in guna so pacifies Pitta. Pitta is the main factor for Kamala. Ghritkumari is yakrita uttejaka and Artavjanana action. So, clearance of srotodushti which is sanga in case of Kamala. Once the srotodushti is cleared, the vicious cycle of provocation of Pitta, is interrupted and relief in symptoms become evident. Kakamachi: Kakamachi is a Rasayana and it pacifies tridosha. Due to its Tikta Katu rasa, it alleviates Kapha and Pitta, because of Guru, Snigdha and Ushna guna, it normalizes Vata dosha. Other properties of Kakamachi are Yakrit uttejaka, Moothrala, Deepana, Pachana, Pittashamaka, Rechaka, Rakta shodhaka, and Tapakrama. Homeostasis of doshas causes clearance of srotodushti which is sanga in case of Kamala. Once the srotodushti is cleared, the vicious cycle of provocation of Pitta is interrupted and relief in symptoms becomes evident. Being Deepana, it is ama Pachana. Retained metabolic wastes can be compared to ama, therefore Kakamachi is supposed to bring down the Serum bilirubin level. From the above description it appears that the drug exerts an effect in breaking the pathogenesis of Kamala. They act as Rasayana, improve Dhatu formation, are Tridoshaghna, Vatashamaka, Pitta sarana and therefore relieve signs and symptoms of Kamala. The Drugs have proved effective in relieving cardinal features of Kamala. As the Drugs have established properties, it may be inferred that the Ghritkumari and Kakamachi are safe and suitable in management of Kamala. Regarding mode of action of Ghritkumari and Kakamachi we have rationally discussed above properties and action which might be responsible to bring changes in sign and symptoms of Kamala. This shows majority of action of the drug are due to Guna prabhava. However, observing the outstanding changes in the condition of patients we have opinion that drug acts certainly by Dravya prabhava also. Therefore we may infer that the action of the Drug in improving the sign and symptoms of Kamala patients by Dravya guna prabhava.
However, it was a pilot study to evaluate the Gunakarmukta and efficacy of Ghritkumari and Kakamachi in Kamala and it was carried out with help of limited resources. We observed a great potential in the drugs (combined Ghritkumari and Kakamachi) to provide relief in patients of Kamala.

**Bibliography**

**Books**

1. Pandeya G. Dravya Guna Vijnana, Krishnadas Academy, Varanasi.
2. Yagi A. Aloe Vera 1st Ed., DHC, Tokyo, 1997
3. Frawley D. & Lad V., The Yoga and Herbs, 1986
5. kyokai,Tokyo, 2002 Samhita

8. Bhavamishra, Bhavaprakash Samhita, Vaidya V.M.Gogte

**Thesis** –


**Bibliography for Kakamachi (solanum nigrum)**

22. Astanga Sangraham.
23. Astanga Nighantu.
24. Ayurvedic harmacology and therapeutic uses of medicinal plants
25. Vaidhya V.M.Gogte
26. Ayurvedic drugs and their plant sources
27. V.V.Shivaranjan, Indira balachandra n.
28. A catalogue of Indian synonyms Modern Sheriff
29. A catalogue of medicinal plants exhibits
31. Charaka Samhita.
32. Chakradatta.
33. Classical uses of medicinal plants 5 Sharma P.V. Prof.
34. Dhanvantari Nighantu 5 Sharma P.V.Prof. & Dr.Guruprasad
35. Dravya guna Hastamalaka.
36. Dravya guna Kosha 5 Sharma P.V.Prof.
37. Dravya guna vijnana Vol.II 5 Sharma P.V.Prof.
38. Flora of Coorg [Karnataka] India 5 S.N.Yoganarasimhan
39. Glossary of vegetable drugs in Brahatrayee 5 Takur Balwant Singh & Dr.K.C.Chunekar 1
40. Hrudayadeepika Nighantu & Siddhamantra 5 Vaidhyacharya Kesava with prakasha commentary of Vopadeva.
41. Haritakyad Nighantu
42. Indian material medica Vol.I 5 Dr.K.M.Nadkarni
43. Indian medicinal plants 5 A compendium of 500 species, Orient Longman.
44. Indian medicinal plants Vol.III 5 K.R. Kirthikar & B.D.Basu
45. Kaiyadeva Nighantu
46. Madapala Nighantu 33
47. Medicinal plants of India Vol.I [Karnataka] 5 S.N.Yoganarasimhan
48. Medicinal plants of India Vol.II [Tamil Nadu]
49. Materia medica of Hindus5 Uday chand dutta
50. Nighantu Adarsha 5 Vaidhya Sri Bapalal
51. Raja Nighantu 5 Tripati Dr.Indradev
52. Sushruta Samhita
53. Shaligrama Nighantu 5 Lal Shastri
54. Sabdakalpa druma
55. Sivakosha of Sivadatta Misra
56. Sarangadara Samhita 5 English commentary
57. Taxonomy of vascular plants 5 George H.M. Lawrence
58. The Wealth of India.
59. Vanoushadi Nidarshika

Bibliography for Kamala roga.
72. Davidson, s Principlaes and practice of Medicine, 20 th editions, 2006.
73. Harison, s principle of internal medicine: 12th Ed.15. Human Anatomy: By B.D.Chaurasia.
74. Sushruta Samhita: Commentary by Dr.Ambika Datta Shastri; Chaukhambha Sanskrita Sansthan, 1972.