

**Evaluation of Hilo<sup>®</sup> versus Daflon<sup>®</sup> in patients  
suffering from hemorrhoids:  
A randomized, controlled, open-labelled,  
multicentric study**

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**ABSTRACT**

**Aims:** To evaluate and compare the efficacy of Hilo<sup>®</sup> and Daflon<sup>®</sup> 500 mg, in the treatment of hemorrhoids.

**Study design:** It is a multicentric, randomized, comparative clinical trial conducted for the period of 15 days.

**Place and Duration of Study:** Janta Hospital and Maternity Centre, Varanasi; King George Memorial Hospital, Lucknow; Vijan Hospital and Research Centre, Nasik and Santosh Hospital, Bangalore between May 2018 and December 2019.

**Methodology:** 201 patients were screened and 200 patients with hemorrhoids (proctoscopy proven Grade I to III) were randomly assigned to receive either Hilo<sup>®</sup> capsules (n = 99) or Daflon<sup>®</sup> 500 mg tablets (n = 101). Assessment of hemorrhoidal symptoms was carried out in all patients on Day 7 and Day 15. Proctoscopic examination was carried out before the start of treatment i.e. on day 0 and at the end of treatment duration i.e. on day 15

**Results:** The patients treated with Hilo<sup>®</sup> showed a statistically significant improvement in the clinical symptoms of bleeding, pain, itching, soiling, tenesmus, irritation after defecation and constipation on day 7 and day 15 as compared to baseline. The "mean total symptom score" reduced by  $4.55 \pm 2.07$  vs  $3.44 \pm 2.00$ ;  $P < .0001$  on day 7 and  $7.56 \pm 2.40$  vs  $6.22 \pm 2.55$ ;  $P < .0001$  on day 15 in the patients treated with Hilo<sup>®</sup> and Daflon<sup>®</sup> respectively. In Hilo<sup>®</sup> Group 82.83% of patients assessed that the treatment with Hilo<sup>®</sup> made them 'A lot better' as compared to only 48.51% in Daflon<sup>®</sup> group. In the Hilo<sup>®</sup> group 20.2% of patients' treatment outcome was assessed as 'Excellent' by the investigators as compared to only 0.99% of patients in Daflon<sup>®</sup> group. No major adverse events were reported in the study with the use of either product.

**Conclusion:** Hilo<sup>®</sup> is found to provide better reduction in clinical symptoms of patients suffering from hemorrhoids as compared to Daflon<sup>®</sup>.

*Keywords: Hemorrhoids, clinical study, phlebotonics, flavonoids, catechins*

**1. INTRODUCTION**

Hemorrhoids are defined as the symptomatic enlargement and distal displacement of the normal anal cushions. The most common symptom of hemorrhoids is rectal bleeding associated with bowel movement. The abnormal dilatation and distortion of the vascular channel, together with destructive changes in the supporting connective tissue within the anal cushion, is a paramount finding of hemorrhoids [1].

24 Approximately 40.7 million people in India are reported to suffer from haemorrhoids [2].  
25 Hemorrhoidal symptoms are observed in about 60% of the patients suffering from  
26 hemorrhoids. The most common symptom of internal hemorrhoids is bleeding which can be  
27 painless and is bright red in color. The external hemorrhoids are more likely to be associated  
28 with pain, due to activation of perianal innervations associated with thrombosis. Patients  
29 typically describe a painful perianal mass that is tender to palpation. The other symptoms of  
30 hemorrhoids include: tenesmus, irritation of the skin surrounding the anus, soiling, itching,  
31 mucus discharge, sensation of tissue prolapse etc [3,4].

32 For the management of hemorrhoids, lifestyle changes and other non-operative measures  
33 have been recommended as first line therapy for management of hemorrhoids. These  
34 measures for hemorrhoids management are associated with significant improvement in the  
35 outcome scores reported by patients [5].

36 Various other options are available which are classified as surgical management such as  
37 Hemorrhoidectomy, Stapled hemorrhoidopexy, Doppler-guided hemorrhoid artery ligation.  
38 Non-surgical office procedures or minimally invasive procedures are also commonly used  
39 like Rubber band ligation, Sclerotherapy, Infrared coagulation, whereas the conservative  
40 management entailing treatment with phlebotonics, consumption of high fibre foods, psyllium  
41 husks, topical creams to relieve inflammation and pain, sitz bath, analgesics etc. is used for  
42 the symptomatic management of the hemorrhoids [3,6].

43 The medical and conservative management with high-fibre diets, stool softeners and  
44 laxatives are the preferred treatments for grade I to grade II hemorrhoids whereas surgical  
45 procedures are reserved for the more severe hemorrhoids [7].

46 Phlebotonics are a heterogeneous group of drugs which are indicated for the treatment of  
47 chronic vein insufficiency and also for the management of less severe hemorrhoids. These  
48 drugs are helpful in the management of Grade-I, Grade-II as well as thrombosed  
49 hemorrhoids. Phlebotonics act by strengthening the vascular walls which increases the  
50 venous tone and improves lymphatic drainage thus normalises the capillary permeability.  
51 Phlebotonics are mostly natural products; e.g.: flavonoids, saponides, etc. The synthetic  
52 phlebotonics include: calcium dobesilate, naftazone, aminaftone, chromocarbe [8].

53 Hilo® capsules are rich in flavonoid contents and act as phlebotonic with vascular-protecting  
54 properties. Hilo® reinforces venous tone, decreases venous capacitance, venous  
55 distensibility and venous emptying time. [9] Hilo® protects the microcirculation by fighting the  
56 venous inflammation via decreasing leukocyte activation, and as a consequence, by  
57 inhibiting the release of inflammatory mediators (Cytokines, IL 1-β & TNFα), free radicals (5-  
58 LOX, ROS & RNS) and prostaglandins. Thus, Hilo® normalizes capillary permeability and  
59 strengthens capillary resistance. [10]

60 Hilo® also acts on the lymphatic system and improves lymphatic drainage by increasing  
61 lymph flow and lymph oncotic pressure. [11] This action on the lymphatic system is  
62 associated with a venotonic and vasculoprotective effect thereby reducing edema. By virtue  
63 of its venotonic, vascular-protecting and anti-inflammatory action, Hilo® improves  
64 hemorrhoidal signs and symptoms e.g., anal discomfort, pain, redness, anal discharge,  
65 tenesmus, pruritus, erythema and bleeding. In addition to the above it also significantly  
66 reduces the frequency, severity and duration of acute hemorrhoidal episodes and bleeding in  
67 all grades of haemorrhoids. [12]

68 Daflon® tablets are made from micronized purified flavonoid fraction consisting of 10%  
69 hesperidin and 90% diosmin. Daflon® also belongs to the phlebotonic category of drugs.

70 Daflon® exerts its effect by increasing the venous tone, protecting the microcirculation  
71 against inflammatory process and improving the lymphatic drainage [10]. It is indicated  
72 clinically for the treatment of venous insufficiency and hemorrhoids [13].

73 The present study was conducted to compare the efficacy and safety of Hilo® and Daflon®,  
74 in the treatment of hemorrhoids.

75

## 76 **2. MATERIAL AND METHODS**

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### 78 **2.1 STUDY DESIGN**

79 This was a multicentric, randomized, open labeled study. All the patients were randomly  
80 assigned in a 1:1 ratio that is 101 patients in Hilo® group and 99 in Daflon® group  
81 respectively. The study was conducted as per the ICH GCP guidelines and Schedule-Y of  
82 Indian drugs and cosmetics act. The respective institutional ethics committees of the trial  
83 sites approved the study protocol and other relevant documents before the enrolment of  
84 patients.

### 85 **2.2 PARTICIPANTS**

86 For inclusion in the study, the patients had to qualify the inclusion and exclusion criteria as  
87 per the approved protocol. Adult patients of either gender diagnosed with hemorrhoids  
88 confirmed by proctoscopy were included in the study. All the eligible patients were provided  
89 with all the necessary information regarding the study and the investigational products and  
90 were asked to sign the informed consent form before proceeding with enrolment in study.

91 Patients using other anti-hemorrhoidal drugs or planning to undergo any surgical procedure  
92 for hemorrhoids and pregnant women, or lactating mothers were not included in the study.

93 This study was conducted at four centers in India- Janta Hospital and Maternity Centre,  
94 Varanasi; King George Memorial Hospital, Lucknow; Vijan Hospital and Research Centre,  
95 Nasik and Santosh Hospital, Bangalore.

### 96 **2.3 INTERVENTIONS**

97 The enrolled patients were randomized to receive either of the two investigational products:  
98 Hilo® herbal capsules (2 caps twice daily), manufactured by Zuventus Healthcare Ltd. India  
99 and Daflon® 500 mg (2 tablets daily), manufactured by Serdia Pharmaceuticals (India) Pvt.  
100 Ltd.

101 Hilo® is a herbal preparation containing a mixture of four herbs, where each capsule  
102 contains Commiphora molmol (Heerabol) oleoresin (250mg), Gardenia gummifera  
103 (Naadihingu) gum-resin (83mg), and Tagates erecta (Genda) flowers (83.5mg), and Mesua  
104 ferrea (Nagakesar) stem (83.5). It is standardized to contain not less than 7% of total  
105 catechins and epicatechins. Each Daflon® 500 mg tablet contains micronized purified  
106 flavonoid extracts of rutaceae 500 mg, equivalent to 450 mg of diosmin, and 50 mg of  
107 hesperidine per tablet.

108 At the end of treatment regimen (Day 15), study medication containers were retrieved from  
109 the patients and the remaining tablets were counted. Thus the compliance was ensured by

110 project staff through pill count. Any other anti-hemorrhoidal treatment or laxatives were not  
111 allowed during the trial period.

## 112 **2.4 OUTCOMES**

113 Primary outcome of the study was to evaluate an improvement in the intensity of  
114 hemorrhoidal symptoms and grades of haemorrhoids as observed with proctoscope on Day  
115 15 by the investigator.

116 Secondary outcome was global assessment for overall improvement by patient and  
117 physician on Day 15 and number of adverse reactions reported by patients throughout the  
118 study duration.

## 119 **2.5 SAMPLE SIZE**

120 A sample size of the study was calculated using a level of significance of 5% and a power of  
121 90%. Following parameters were considered-  
122 Confidence level = 95%  
123 Acceptable difference = 0.10  
124 Assumed proportion = 0.50  
125 Using WINPEPI software, it was found that required sample size is 97 in each group.  
126 The present study enrolled 200 patients, 100 patients in each group.

## 127 **2.6 RANDOMISATION**

128 Computer generated randomization blocks were generated to assign the eligible patients to  
129 either of the treatment groups. Patients were screened and enrolled by the investigators  
130 based upon the eligibility criteria. After obtaining written informed consent, Hilo® herbal  
131 capsules or Daflon® 500 mg tablet was dispensed to each patient by investigators as per the  
132 randomization chart.

## 133 **2.7 STUDY ASSESSMENTS**

134 On day 0, screening and randomization of the patients was done. This involved signing of  
135 informed consent document and enrolment of patients as per the inclusion and exclusion  
136 criteria. Demographics and medical history including previous history of hemorrhoids or any  
137 other chronic diseases was assessed. The investigational products were dispensed to all  
138 eligible patients on day 0 as per the randomization scheme generated using SAS 9.1  
139 software. The enrolled patients were given the investigational products for 15 days with  
140 instructions for drug administration.

141 Proctologic examination was performed to assess the hemorrhoidal conditions on day 0 and  
142 day 15 i.e. before the start of treatment and at the end of the study. Proctologic assessment  
143 was performed in the left-lateral position by inspection of the anal verge of the anal canal by  
144 using a proctoscope. Parameters namely, grade (I, II, III, IV) and position of hemorrhoids (at  
145 one site, two sites or all three primary sites, i.e., 3'O clock, 7'O clock, 11'O clock position)  
146 were assessed. The severity of clinical symptoms of hemorrhoids (bleeding, pain, itching,  
147 soiling, tenesmus, irritation after defecation and constipation) was assessed using a 4-point  
148 scale: (0= absent, 1= mild, 2= moderate and 3= severe) on each visit i.e. day 0, day 7 and  
149 day 15 Additionally on day 15, global assessment of the interventions was done subjectively  
150 by the patients as well as the investigator.

151 **2.8 STATISTICAL ASSESMENT**

152 The following null hypothesis was formulated:

153 Ho: There is no difference between the two treatment groups in improving the clinical  
154 symptoms of hemorrhoids

155 H1: There is a difference between the two treatment groups in improving the clinical  
156 symptoms of hemorrhoids

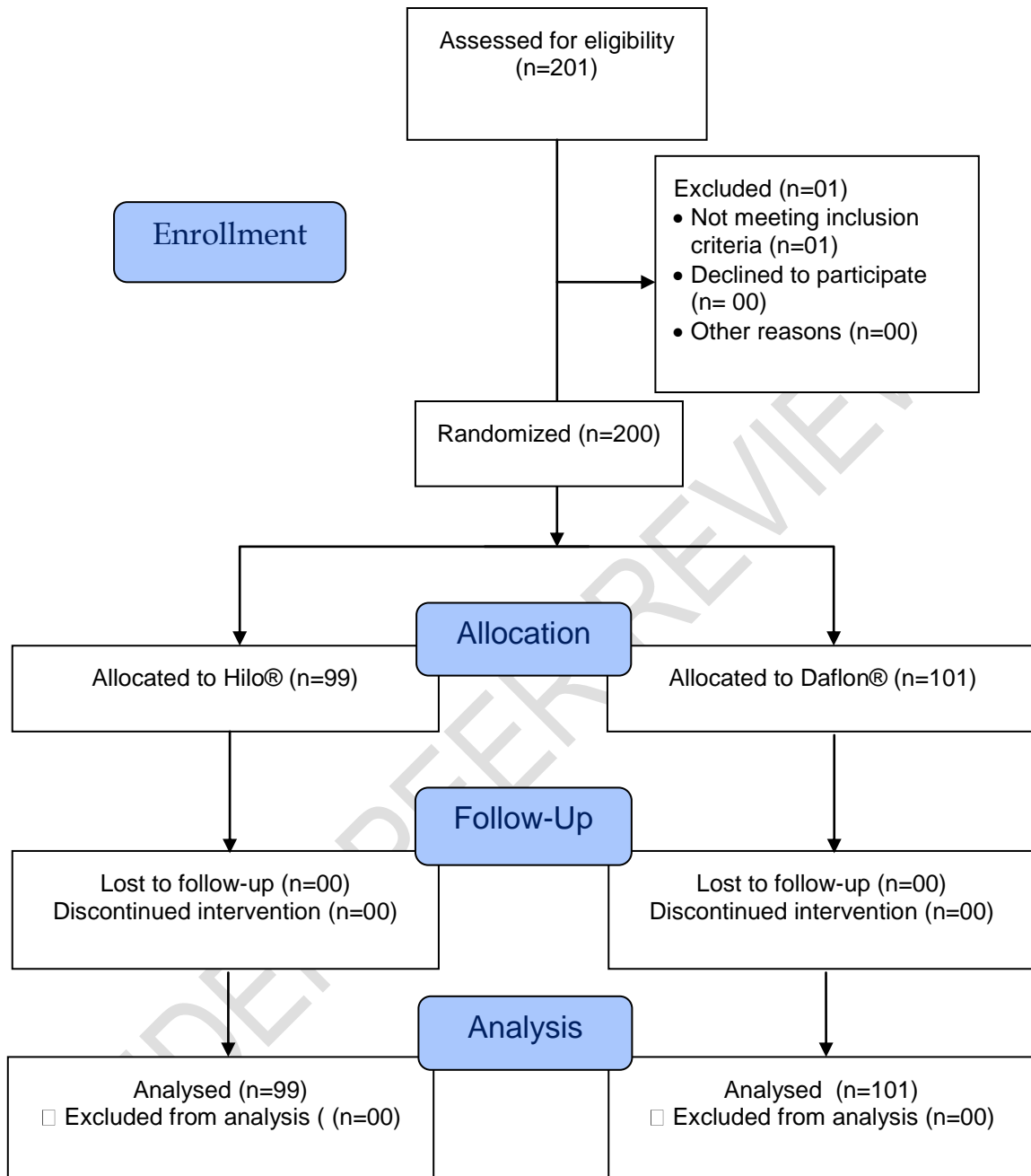
157 Analysis were performed using StatsDirect software (Version 3.1.22). The data are  
158 expressed as mean  $\pm$  SD or percentage. Unpaired t-test was used to compare the  
159 demographic parameters of age, weight and height. Mann-Whitney U test was used for  
160 'between the group' comparison while Wilcoxon's signed ranks test was used to compare  
161 the changes 'within the group' and McNemar and exact (Liddell) test was used to compare  
162 the proportions.

163 95% Confidence Intervals (CIs) for the true proportions were also calculated. All 'P values'  
164 were considered significant if less than .05.

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166 **3. RESULTS**

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168 The discussion should not repeat the results, but provide detailed interpretation of data. This  
169 should interpret the significance of the findings of the work. Citations should be given in  
170 support of the findings. The results and discussion part can also be described as separate, if  
171 appropriate.

172  
173 In the present study, a total of 201 patients presenting with hemorrhoids were screened at 4  
174 clinical trial sites. Out of the 201 screened, 200 patients fulfilled the eligibility criteria and  
175 were enrolled in the study. The patients were randomised to receive either Hilo® or Daflon®  
176 where the Hilo® group comprised of 99 patients while 101 patients were allotted in the  
177 Daflon® group. All 200 patients completed the study as per the approved protocol and their  
178 data was subjected to statistical analysis at the end of the study. A consort chart of trial  
179 participation is described in Figure 1. The first patient was enrolled on 18<sup>th</sup> May 2017 at  
180 Janta Hospital, Varanasi and the last patient completed the study on 28<sup>th</sup> December 2018 at  
181 Santosh Hospital, Bangalore.



182

183 **Fig. 1. Consort chart of trial participants.**

184 *n= number of patients*

185 **3.1 DEMOGRAPHICS**

186 On Day 0, the demographic parameters like age, height and weight were documented. The  
 187 means of demographic parameters of age, weight and height were compared using unpaired  
 188 t-test. The baseline individual symptom scores of the two treatment groups were compared  
 189 using Mann-Whitney U test. No significant difference was observed between the two groups  
 190 ( $P > .05$ ) (Table 1) at baseline keeping the randomization unbiased.

191 **Table 1. Comparative demographics using unpaired t-test and baseline scores of**  
 192 **clinical symptoms using Mann-Whitney U Test**

	<b>Hilo®</b> (n=99) Mean ± S.D.	<b>Daflon®</b> (n=101) Mean ± S.D.	<b>P</b>	<b>95% C.I.</b>
<b>Age (years)</b>	40.17 ± 13.67	38.31 ± 12.51	.32	Mean diff= 1.86 -1.79 to 5.52
<b>Weight (kg)</b>	61.07 ± 10.87	60.38 ± 8.29	.18	Mean diff= 0.69 0.34 to 0.53
<b>Height (cm)</b>	161.83 ± 7.51	161.46 ± 7.45	.61	Mean diff= 0.37 -1.71 to 2.46
<b>Baseline scores of clinical symptoms</b>				
<b>Bleeding</b>	1.45 ± 0.52	1.57 ± 0.64	.24	Mean diff= -0.12 0.41 to 0.57
<b>Pain</b>	1.75 ± 0.52	1.72 ± 0.51	.73	Mean diff= 0.03 0.46 to 0.62
<b>Itching</b>	1.33 ± 0.64	1.29 ± 0.67	.83	Mean diff= 0.04 0.41 to 0.57
<b>Soiling</b>	1.27 ± 0.62	1.33 ± 0.60	.63	Mean diff= -0.06 0.44 to 0.59
<b>Tenesmus</b>	1.54 ± 0.63	1.51 ± 0.66	.99	Mean diff= 0.03 0.42 to 0.58
<b>Irritation while defecation</b>	1.24 ± 0.70	1.38 ± 0.75	.14	Mean diff= -0.14 0.47 to 0.63
<b>Constipation</b>	1.79 ± 0.67	1.71 ± 0.60	.33	Mean diff= 0.08 0.39 to 0.55

193 \* Mean ± S.D.= Mean ± Standard Deviation, C.I.= Confidence Interval, kg = kilogram, cm= centimeter

194 **3.2 Assessment of Total Symptom Score and Individual Clinical Symptom Score**

195 The total symptom score was calculated by adding the individual symptom scores of  
 196 bleeding, pain, itching, soiling, tenesmus, irritation after defecation, constipation for each  
 197 patient. The individual symptom scores of bleeding, pain, itching, soiling, tenesmus, irritation  
 198 after defecation, constipation were scored for their severity on Day 0, Day 7 and Day 15. The  
 199 mean change in the total symptom score and the individual clinical symptom score from the  
 200 baseline score of Day 0 was evaluated at Day 7 and Day 15 using Wilcoxon's signed ranks  
 201 test for both the groups (Table 2).  
 202

203 **Table 2 Improvement in the total symptom score and individual clinical symptom**  
 204 **score of hemorrhoids before and after treatment with Hilo® and Daflon® using**  
 205 **Wilcoxon's signed ranks test**

	Hilo® (n=99)					Daflon® (n=101)				
	Day 0 (Visit 1) Mean ± S.D.	Day 7 (Visit 2) Mean ± S.D.	Day 15 (Visit 3) Mean ± S.D.	Mean Change from Day 0 to Day 7 (95% C.I.)	Mean Change from Day 0 to Day 15 (95% C.I.)	Day 0 (Visit 1) Mean ± S.D.	Day 7 (Visit 2) Mean ± S.D.	Day 15 (Visit 3) Mean ± S.D.	Mean Change from Day 0 to Day 7 (95% C.I.)	Mean Change from Day 0 to Day 15 (95% C.I.)
<b>Bleeding</b>	1.45 ± 0.73	0.69 ± 0.62	0.23 ± 0.47	0.77±0.65* (0.5 to 1)	1.22±0.72* (1.0 to 1.5)	1.57 ± 0.64	0.99 ± 0.56	0.43 ± 0.65	0.58±0.55* (0.5 to 0.5)	1.15±0.80* (0.99 to 1.31)
<b>Pain</b>	1.75 ± 0.52	0.94 ± 0.47	0.68 ± 0.68	0.81±0.65* (0.5 to 1)	1.07±0.92* (1.0 to 1.5)	1.72 ± 0.51	1.14 ± 0.58	0.86 ± 0.65	0.58±0.53* (0.5 to 0.5)	0.86±0.82* (0.5 to 1)
<b>Itching</b>	1.33 ± 0.64	0.89 ± 0.50	0.64 ± 0.61	0.43±0.66* (0.5 to 0.5)	0.69±0.91* (0.5 to 1)	1.29 ± 0.67	1.00 ± 0.57	0.80 ± 0.55	0.29±0.61* (0.0 to 0.5)	0.49±0.74* (0.5 to 0.5)
<b>Soiling</b>	1.27 ± 0.62	0.54 ± 0.56	0.21 ± 0.41	0.74±0.58* (0.5 to 1)	1.06±0.62* (1.0 to 1.0)	1.33 ± 0.60	0.76 ± 0.59	0.39 ± 0.53	0.56±0.59* (0.5 to 0.5)	0.94±0.61* (1.0 to 1.0)

	Hilo® (n=99)					Daflon® (n=101)				
	Day 0 (Visit 1) Mean ± S.D.	Day 7 (Visit 2) Mean ± S.D.	Day 15 (Visit 3) Mean ± S.D.	Mean Change from Day 0 to Day 7 (95% C.I.)	Mean Change from Day 0 to Day 15 (95% C.I.)	Day 0 (Visit 1) Mean ± S.D.	Day 7 (Visit 2) Mean ± S.D.	Day 15 (Visit 3) Mean ± S.D.	Mean Change from Day 0 to Day 7 (95% C.I.)	Mean Change from Day 0 to Day 15 (95% C.I.)
<b>Tenesmus</b>	1.54 ± 0.63	0.84 ± 0.63	0.75 ± 0.68	0.69±0.54* (0.5 to 1)	0.79±0.81* (0.5 to 1)	1.51 ± 0.66	0.99 ± 0.62	0.90 ± 0.61	0.52±0.58* (0.5 to 0.5)	0.61±0.71* (0.5 to 1)
<b>Irritation after defecation</b>	1.24 ± 0.70	0.81 ± 0.55	0.56 ± 0.59	0.43±0.76* (0.28 to 0.59)	0.69±0.85* (0.5 to 1)	1.38 ± 0.73	1.00 ± 0.65	0.79 ± 0.62	0.39±0.75* (0.5 to 0.5)	0.59±0.74* (0.5 to 1)
<b>Constipation</b>	1.79 ± 0.67	1.06 ± 0.53	0.66 ± 0.48	0.73±0.53* (0.5 to 1)	1.13±0.69 (1.0 to 1.5)	1.71 ± 0.60	1.18 ± 0.65	0.82 ± 0.49	0.53±0.61* (0.5 to 0.5)	0.89±0.66* (1.0 to 1.0)
<b>TOTAL SYMPTOM SCORE</b>	10.75 ± 2.02	6.20 ± 1.71	3.19 ± 1.54	4.55±2.07* (4.5 to 5)	7.56±2.40* (7 to 8)	10.70 ± 2.40	7.26 ± 2.19	4.48 ± 2.05	3.44±2.00* (3 to 4)	6.22±2.55* (5.5 to 7)

206  $P < .0001$

207 Mean changes in individual symptom scores from baseline to day 7 were statistically  
208 significantly improved in Hilo® group as compared to Daflon® group viz. Bleeding (0.77 ±  
209 0.65 vs 0.58 ± 0.55;  $P < .0001$ ), Pain (0.81 ± 0.65 vs 0.58 ± 0.53;  $P < .0001$ ), Itching (0.43 ±  
210 0.66 vs 0.29 ± 0.61;  $P < .0001$ ), Soiling (0.74 ± 0.58 vs 0.56 ± 0.59;  $P < .0001$ ), Tenesmus  
211 (0.69 ± 0.54 vs 0.52 ± 0.58;  $P < .0001$ ), Irritation after defecation (0.43 ± 0.76 vs 0.39 ± 0.75;  
212  $P < .0001$ ) and Constipation (0.73 ± 0.53 vs 1.13 ± 0.69;  $P < .0001$ ).

213 Mean changes in individual symptom scores from baseline to day 15 were statistically  
214 significantly improved in Hilo® group as compared to Daflon® group viz. Bleeding (1.22 ±  
215 0.72 vs 1.15 ± 0.80;  $P < .0001$ ), Pain (1.07 ± 0.92 vs 0.86 ± 0.82;  $P < .0001$ ), Itching (0.69 ±  
216 0.91 vs 0.49 ± 0.74;  $P < .0001$ ), Soiling (1.06 ± 0.62 vs 0.94 ± 0.61;  $P < .0001$ ), Tenesmus

217 (0.79 ± 0.81 vs 0.61 ± 0.71;  $P < .0001$ ), Irritation after defecation (0.69 ± 0.85 vs 0.59 ± 0.74;  
 218  $P < .0001$ ) and Constipation (1.13 ± 0.69 vs 0.89 ± 0.66;  $P < .0001$ ).  
 219 Mean change in total symptom score was found to be statistically significant from baseline to  
 220 day 7 (4.55 ± 2.07 vs 3.44 ± 2.00;  $P < .0001$ ) and day 15 (7.56 ± 2.40 vs 6.22 ± 2.55;  $P <$   
 221  $.0001$ ) in Hilo® group as compared to Daflon® group.

222 The patients treated with Hilo® capsules showed a significantly better improvement in total  
 223 symptom score on both Day 7 & Day 15 when compared to Daflon® ( $P < .0001$ ) (Table 3).

224 **Table 3 Difference between Hilo® and Daflon® treatment groups in improvement of**  
 225 **total symptom score of hemorrhoids using Mann-Whitney U test**

Improvement in Total Symptom Score with Hilo® and Daflon®	
Improvement on Day 7	Mean diff = 1.10 95% C.I. = 0.27 to 0.41
Improvement on Day 15	Mean diff = 0.23 95% C.I. = 0.04 to 0.11

226  $P < .0001$

227 **3.3 Proportion of patients exhibiting reduction in total symptom score**

228 The proportion of patients showing reduction of ≥ 50%, ≥ 75% and ≥ 80% in the total  
 229 symptom score were evaluated and the two groups were compared using McNemar and  
 230 exact (Liddell) test (Table 4).

231 In Hilo® treatment group, the scores of 89.89% patients (89/99) was reduced to ≥ 50% on  
 232 day 15. In the Daflon® treatment group 74.26% patients (75/101) exhibited ≥ 50% reductions  
 233 in total symptom score on Day 15. 48.48% patients from Hilo® group while only 16.83%  
 234 patients receiving Daflon® achieved ≥ 75% reduction in total symptom score. The total  
 235 symptom score of 32.32% patients from Hilo® group and 13.86% patients from Daflon®  
 236 group improved by ≥ 80% on Day 15.

237 The number of patients achieving ≥ 50%, ≥ 75% and ≥ 80% reduction in total symptom score  
 238 on Day 15 was significantly higher in the Hilo® group when compared to Daflon® group ( $P <$   
 239  $.0001$ ).

240 **Table 4. Number of patients showing ≥ 50%, ≥ 75% and ≥ 80% reduction in total**  
 241 **symptom score on Day 15 as compared to Day 0 using McNemar and exact (Liddell)**  
 242 **test**

Improvement in hemorrhoid symptoms	Hilo® group (n= 99)	Daflon® group (n=101)	P value
Number of patients with ≥ 50% reduction in total symptom score on	89	75	< .0001

Number of patients with $\geq$ 75% reduction in total symptom score on	48	17	< .0001
Number of patients with $\geq$ 80% reduction in total symptom score on	32	14	< .0001

243

244 **3.4 Presence of clinical symptoms before and at the end of treatment period**

245 At the baseline, the number of patients exhibiting the various clinical symptoms of  
 246 hemorrhoids (bleeding, pain, itching, soiling, tenesmus, irritation after defecation, and  
 247 constipation) were identified. At the end of the treatment i.e. Day 15, the proportion of  
 248 patients exhibiting the presence of these clinical symptoms were evaluated. The proportion  
 249 of patients exhibiting clinical symptoms in the two groups were compared at the end of the  
 250 treatment using McNemar and exact (Liddell) test (Table 5).

251 On comparing the two treatment groups, it was noted that a significantly less ( $P < 0.01$ )  
 252 number of patients from Hilo® group exhibited the clinical symptoms of bleeding, pain,  
 253 itching, soiling, tenesmus, irritation after defecation and constipation as compared to  
 254 Daflon® on Day 7 as well as Day 15.

255 **Table 5 Patients exhibiting clinical symptoms of hemorrhoids at baseline, day 7 and**  
 256 **day 15 using McNemar and exact (Liddell) test**

Clinical symptoms	Hilo® (n=99)			Daflon® (n=101)			Hilo® vs. Daflon® P value	
	Number of patients exhibiting clinical symptoms			Number of patients exhibiting clinical symptoms			Comparison of proportion of patients exhibiting clinical symptoms	
	Day 0	Day 7	Day 15	Day 0	Day 7	Day 15	Day 7	Day 15
Bleeding	87	61	21	94	86	34	< .0001	< .01
Pain	97	85	55	99	90	72	< .0001	< .01
Itching	91	81	56	89	85	68	< .0001	< .01
Soiling	90	50	21	95	69	37	< .01	< .01
Tenesmus	95	70	62	94	81	77	< .0001	< .0001
Irritation after	85	73	50	88	80	69	< .0001	< .01

<b>defecation</b>								
<b>Constipation</b>	96	88	65	100	89	78	< .0001	< .0001

257

258 **3.5 Improvement in Hemorrhoidal Grades**

259 Hemorrhoidal assessment was performed as described in the study assessment section. At  
 260 the end of study (Day 15), the improvement in the stage of hemorrhoids was significantly  
 261 higher in the Hilo® group (Grade I: 74.75% of patients) as compared to Daflon® group  
 262 (Grade I: 55.45% of patients) (Table 6).

263 **Table 6 Proportion of patients exhibiting various grades of hemorrhoids on**  
 264 **proctoscopic examination on Day 0 and Day 15**

	<b>Hilo® (n=99)</b>		<b>Daflon® (n=101)</b>	
	Day 0 n (%)	Day 15 n (%)	Day 0 n (%)	Day 15 n (%)
<b>Grade IV</b>	00 (0%)	00 (0%)	01 (0.99%)	00 (0%)
<b>Grade III</b>	16 (16.16%)	02 (2.02%)	25 (24.75%)	01 (0.99%)
<b>Grade II</b>	67 (67.68%)	21 (21.21%)	59 (58.42%)	43 (42.57%)
<b>Grade I</b>	16 (16.16%)	74 (74.75%)	16 (15.84%)	56 (55.45%)
<b>No Hemorrhoids</b>	00 (0%)	02 (2.02%)	00 (0%)	01 (0.99%)

265

266 At the end of the study, with Hilo® treatment 62 patients out of 83 (74.69%) from baseline of  
 267 Grade II and Grade III combined exhibited improvement to Grade I. Similarly, with Daflon®  
 268 treatment 44 patients out of 86 (total of Grade II and Grade III) improved to Grade I  
 269 (51.16%). It was observed that a significantly a greater number of patients ( $p < 0.01$ ) from  
 270 Hilo® group showed improvement in hemorrhoidal grade as compared to Daflon® (Table 7).  
 271 This is an important aspect, as Grade III hemorrhoids, unlike Grade I and II hemorrhoids, do  
 272 not usually present spontaneous improvement of the symptoms.

273 **Table 7 Difference between Hilo® and Daflon®: Proportion of patients showing**  
 274 **improvement from Grade III & II (combined) III to Grade I using McNemar and exact**  
 275 **(Liddell) test**

	Visit 1 (Baseline)	Visit 3 (Day 15)	Success Rate	Hilo® vs. Daflon®
	Grade III & II (n)	Grade I (n)		P value
Hilo®	83	62	74.69 %	.006*
Daflon®	86	44	51.16 %	

276 P < .01

277 **3.6 Global Assessment of Therapy (Table 8)**

278 In Hilo® Group 82.83% of patients assessed that the treatment with Hilo® made them 'A lot  
279 better' as compared to only 48.51% in Daflon® group.

280 In the Hilo® group 20.2% of patients' treatment outcome was assessed as 'Excellent' by the  
281 investigators while in Daflon® group only 0.99% patients showed 'Excellent' outcome as per  
282 the investigator. 5.94% of patients in Daflon® group showed 'Poor' outcome at the end of  
283 study.

284 **Table 8 Assessment of therapy by patients and investigators**

Assessment Of Therapy	Hilo® Group (n=99)	Daflon® Group (n=101)
<b>By Patients</b>		
The treatment made me a lot worse	0 (0%)	0 (0%)
The treatment made me slightly worse	0 (0%)	0 (0%)
The treatment made no change to my symptoms	01 ( 1.01% )	9 ( 8.91% )
The treatment made me slightly better	15 ( 15.15% )	43 (42.57%)
The treatment made me a lot better	82 (82.83% )	49 (48.51%)
The treatment completely relieved my symptoms	01 (1.01% )	0 (0% )
<b>By Investigators</b>		
Excellent	20 (20.20%)	1 (0.99%)
Good	67 (67.68% )	59 (58.42% )
Satisfactory	11 (11.11%)	35 (34.65%)
Poor	0 (0% )	6 (5.94% )

285

### 286 **3.7 Adverse Events**

287 There were no adverse events reported/observed in patients of either treatment groups  
288 during the course of the study.

## 289 **4. DISCUSSION**

290 A major component of a safe and effective therapy for hemorrhoids is the use of herbal  
291 products. Several herbal extracts containing flavonoids have been shown to improve  
292 microcirculation, capillary flow, and vascular tone, and strengthen connective tissue of the  
293 perivascular amorphous substrate. Flavonoid molecules also reduce inflammation by  
294 inhibiting prostaglandin and free radicals generated during the inflammatory response. The  
295 standard treatments for hemorrhoids are aimed toward removing the problem or palliating  
296 the disease.

297 Additionally, the low compliance associated with treatments such as hydrotherapy,  
298 mechanical compression therapy, and diet and lifestyle changes. This renders oral dietary  
299 supplementation an attractive option. The use of oral flavonoids offers an effective approach  
300 for the treatment of hemorrhoids. Early intervention with conservative therapies may prevent  
301 time-consuming and expensive complications of haemorrhoids <sup>[14,15]</sup>.

302 Flavonoids are considered as phlebotonics and were first described in the treatment of  
303 chronic venous insufficiency and edema. They appeared to be capable of increasing  
304 vascular tone, reducing venous capacity, decreasing capillary permeability, and facilitating  
305 lymphatic drainage as well as having anti-inflammatory effects <sup>[1]</sup>.

306 In an earlier study comparing Roidosanal® (standardized to contain not less than 7% of total  
307 catechins and epicatechins) and Daflon®, it was found that both the treatments are equally  
308 effective in improving anorectal conditions and the associated hemorrhoidal clinical  
309 symptoms. No major adverse events were reported in the study with the use of either  
310 product <sup>[16]</sup>.

311 Daflon® tablets are a commercially available herbal medicine in India belonging to a similar  
312 category as that of Hilo®, hence it was used as comparator. In the present study, it was  
313 observed that patients treated with Hilo® showed a significant improvement in the clinical  
314 symptoms of bleeding, pain, itching, soiling, tenesmus, irritation after defecation and  
315 constipation ( $p < 0.0001$ ) on day 7 as well as day 15 as compared to baseline. This is one of  
316 the most important aspect in the treatment of haemorrhoids when patient starts finding  
317 improvement in symptoms, the compliance towards prescribed drug increases and patient  
318 completes the full course of the medicine.

319 The mean total symptom score reduced by 4.55 on day 7 and by 7.56 on day 15 in the  
320 patients treated with Hilo®. The patients treated with Daflon® also showed a significant  
321 reduction in individual symptom score. The mean total symptom score reduced by 3.44 on  
322 day 7 and by 6.22 on day 15 in the patients treated with Daflon®. The improvement in total  
323 symptom score by both the treatments was compared using Mann-Whitney U test. These  
324 symptomatic improvement was corroborated by the proctoscopic findings of reduction in  
325 Grade of haemorrhoids.

326 Hilo® was found to be better in reducing the total symptom score on day 7 and day 15 as  
327 compared to Daflon® ( $p < 0.0001$ ). These results are consistent with previously published

328 data <sup>16</sup>. Thus a 15 days course of Hilo can be recommended before proceeding for  
329 hemorrhoidectomy.

330 The current study has limitation of its smaller sample size. Further studies should be  
331 conducted to observe the impact of 15 days Hilo® therapy in avoiding the surgical  
332 intervention for the treatment of haemorrhoids.

## 333 **5. CONCLUSION**

334 In the present study, Hilo® was found to be better in improving the clinical symptoms of  
335 hemorrhoids as compared to Daflon®. Patients treated with Hilo® also showed improvement  
336 in the grades of hemorrhoids. There were no adverse events reported with either of the  
337 treatments. Hilo® is a safe and effective treatment for hemorrhoids.

338

## 339 **COMPETING INTERESTS DISCLAIMER:**

340 **Authors have declared that no competing interests exist. The products used**  
341 **for this research are commonly and predominantly use products in our area of**  
342 **research and country. There is absolutely no conflict of interest between the**  
343 **authors and producers of the products because we do not intend to use these**  
344 **products as an avenue for any litigation but for the advancement of**  
345 **knowledge. Also, the research was not funded by the producing company**  
346 **rather it was funded by personal efforts of the authors.**

## 347 **COMPETING INTERESTS**

348  
349 Authors have declared that no competing interests exist.

350

## 351 **CONSENT**

352

353 Authors declare that written informed consent was obtained from all the patients who  
354 participated in this study. A copy of the written consent is available for review by the Editorial  
355 office/Chief Editor/Editorial Board members of this journal.

## 356 **ETHICAL APPROVAL**

357

358 All authors hereby declare that all experiments have been examined and approved by the  
359 appropriate ethics committee and have therefore been performed in accordance with the  
360 ethical standards laid down in the 1964 Declaration of Helsinki.

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