ABSTRACT

**Background:** Finasteride is a widely used drug in dermatology for the treatment of androgenetic alopecia. There are many reports of associated sexual side effects. This article reviews the use of once-daily 1 mg finasteride in androgenetic alopecia and its associated sexual adverse effects. **Methods:** A literature search was performed to collect data on the use of finasteride in male pattern baldness. Relevant literature published till March 2014 was obtained from MEDLINE, EMBASE, CINAHL, Cochrane registers and LILACS. The keywords “finasteride”, “male pattern baldness” and “androgenetic alopecia” were used for literature search. Similarly, a search was done for finasteride in female pattern hair loss with keywords “female pattern baldness”, “finasteride” and “female pattern alopecia”. All systematic reviews, meta-analyses, national guidelines, randomized controlled trials, prospective open label studies and retrospective case series in the English literature were reviewed. **Results:** Two hundred sixty two studies were evaluated, twelve of which fulfilled the inclusion criteria. **Conclusions and Recommendations:** Current evidence on the safety of finasteride indicates that it is safe but there is growing concern about its sexual side effects. In view of this, proper information should be provided to patients prior to starting treatment (Level of recommendation 1+, Grade of recommendation B). The reported sexual side effects are few and reverse with stoppage of the drug (Grade of recommendation B) but further studies are required.

**Key words:** Androgenetic alopecia, finasteride, guidelines, sexual side effects

INTRODUCTION

Finasteride is widely used for the treatment of androgen-dependent hair disorders such as androgenetic alopecia.[1] It is a selective 5 alpha reductase inhibitor and is administered orally in a dose of 1 mg once daily for androgenetic alopecia. The bioavailability following oral intake of 1 mg ranges from 26% to 70% with a mean of 65%. It is not related to food intake. The average peak plasma concentration is found to be 9.2 ng/ml. The terminal half-life is approximately five to six hours in men between 18 and 60 years of age and eight hours in men older than 70 years of age.[2] It is extensively metabolized in the liver by cytochrome P450 3A4 enzyme subfamily and excreted in urine and feces. Finasteride has been tried in several doses ranging from 0.2 mg to 5 mg but 1 mg per day is the optimal dose for the treatment of men with male pattern hair loss. There is no difference in efficacy between doses of 1 mg and 5 mg.[3] Long-term daily finasteride is advocated and leads to sustained improvement.[4-6]

METHODS

A literature search was done to collect data on the use of finasteride in male pattern baldness. Relevant literature published until March 2014 was obtained from MEDLINE, EMBASE, CINAHL, LILACS and
Cochrane registers. The keywords “finasteride,” “male pattern baldness” and “androgenetic alopecia” were used. A similar search was done for finasteride in female pattern hair loss with the keywords “female pattern baldness,” “finasteride” and “female pattern alopecia.” All systematic reviews, meta-analyses, national guidelines, randomized controlled trials, prospective open-label studies and retrospective case series were reviewed in the English literature. The available evidence was evaluated using guidelines suggested by the British Association of Dermatologists. The methodology of each study selected was assessed as per the National Institute of Clinical Excellence Technical Manual and graded as “++,” “+” or “−,” based on the extent to which potential biases were minimized. Therefore, a grade of recommendation was assigned as follows: Grade A, at least one meta-analysis, systematic review, or randomized controlled trial rated as 1++, and directly applicable to the target population, or a systematic review of randomized controlled trials or a body of evidence consisting principally of studies rated as 1+, directly applicable to the target population and demonstrating overall consistency of results; Grade B, a body of evidence including studies rated as 2++, directly applicable to the target population and demonstrating overall consistency of results, or extrapolated evidence from studies rated as 1++ or 1++; Grade C, a body of evidence including studies rated as 2+, directly applicable to the target population and demonstrating overall consistency of results, or extrapolated evidence from studies rated as 2++: Grade D, evidence level 3 or 4, or extrapolated evidence from studies rated as 2+, or formal consensus.

RESULTS

Two hundred and sixty two studies were evaluated, twelve of which fulfilled the inclusion criteria for the guidelines.

Efficacy of the drug in androgenetic alopecia

Several publications have established the usefulness of this drug in the management of androgenetic alopecia. A systematic review of twelve studies showed moderate quality evidence that the daily use of oral finasteride increases hair count. It also improves the patient and investigator assessment of hair appearance (evidence level 1+). Long-term use for up to five years has been shown to decrease the likelihood of developing further visible hair loss (evidence level 1+). A study of 270 men with high levels of serum 5 alpha-dihydrotestosterone showed that starting the drug in younger patients had a better response (evidence level 1+). Another study assessed its efficacy over a ten year period and found that efficacy was not reduced with time. In fact, a large proportion of subjects who had no change after a year improved later on maintaining a positive trend with long-term use (evidence level 1++). An Indian study of 100 patients suggested that finasteride alone, or in combination with minoxidil or ketoconazole, showed statistically significant improvement ($P < 0.05$) over minoxidil only recipients (evidence level 1+). The study concluded that a combination of drugs enhanced the efficacy.

A randomized, double-blind, comparative study to determine the efficacy and safety of 3% minoxidil versus combined 3% minoxidil/ and 0.1% finasteride in male pattern hair loss showed significantly greater improvement in the 3% minoxidil/ and 0.1% finasteride group than the minoxidil group (evidence level 1+). Some recent studies with topical finasteride formulations have been published. A double-blind study showed that the therapeutic effects of 1% finasteride gel applied twice daily and oral finasteride, 1 mg daily were relatively similar (evidence level 1+). Topical finasteride can be considered for hair density maintenance after initial improvement with oral finasteride, thereby avoiding the need for using oral finasteride indefinitely.

These studies show that the drug is efficacious in once daily doses of 1 mg. It works better when started early, and the effect is sustained with long term use of up to 10 years. It acts better in combination with other drugs such as minoxidil (grade of recommendation B). Although there was 1 + level of evidence in support of safety, the authors mark it as level ‘B’ of evidence. This is because of a recent publication that claimed adverse side effects could persist even after discontinuation of therapy resulting in the so called post-finasteride syndrome.

Evidence for topical formulations of finasteride is scanty. Only one randomized study was available, therefore it needs more systematic documentation in further studies (grade of recommendation B).

Side effects

The major drawback of finasteride has been side effects on sexual function. This is the major deterrent for patients taking the drug and is responsible for poor compliance. This has received a lot of attention on the internet and in
the press and hence requires proper consideration. There are conflicting reports about the side effects and lack of side effects which are discussed below.

**Studies which found the side effects to be not significant [Table 1]**

Many studies have concluded that the side effects are not significant.[17-22] These studies reveal that sexual adverse effects occur at the rate of 2.1–3.8% (comparable to placebo). Erectile dysfunction is the most common side effect followed by ejaculatory dysfunction and loss of libido. These effects occurred early in therapy and returned to normal on stoppage of the drug or while continuing use of the drug over a period of time. The nocebo effect may explain side effects in some patients.[22] These studies have concluded that the effect of finasteride on sexual functioning is minimal for most men and should not impact the decision to prescribe or take finasteride. A recent review on the available literature arrived at similar conclusions (grade of recommendation B).[23]

| Study                  | Type of study/number of patients | Results                                                                 | Inference                                                                                              | Level of evidence |
|------------------------|---------------------------------|------------------------------------------------------------------------|--------------------------------------------------------------------------------------------------------|-------------------|
| Carbone et al.[17]     | Review of many RCTs - 73 papers | Problems of ejaculation (2.1-7.7%), erection (4.9-15.8%) and decreased libido (3.1-5.4%) | The population studied is different and hence generalization is not possible. So in spite of this being a review of RCTs, we grade it as low evidence | 1+                |
| Mondaini et al.[18]    | 120 patients                    | The incidence of erectile dysfunction, decreased libido and ejaculation disorders were 9.6%, 7.7% and 5.7% for group 1 (5 mg finasteride without counseling) and 30.9%, 23.6% and 16.3% for group 2 (5 mg finasteride with counseling) | There was a significantly higher proportion of sexual dysfunction in patients informed about the sexual side effects (group 2) as compared to those who were not informed | 1+                |
| Kaufman et al.[19]     | 1553 patients                   | Decreased libido, erectile dysfunction and ejaculatory disorders occurred in less than 2% of men. The incidence of each side effect mentioned decreased to ≤0.3% by the 5th year of treatment with finasteride | Incidence of side effects was comparable to that of placebo, both at 1 year and at the end of 5 years | 1+                |
| Leyden et al.[20]      | Multicentric, double-blind, placebo-controlled study done at 45 locations followed by a 1-year open extension | Significant increase in hair count in the frontal scalp of finasteride-treated patients. Efficacy was maintained or improved throughout the 2nd year of the study. Sexual adverse effects were reported in approximately 2% of men in both treatment groups and they resolved while continuing in the study |                                                                                                          | 1+                |
| Moinpour et al.[21]    | Prostate cancer prevention trial. Randomized, double-blind, placebo-controlled, n=18,882 men | Finasteride marginally increased sexual dysfunction and its impact diminished over time; the increase in the sexual activity scale score relative to placebo was 3.21 points (95% CI=2.83-3.59 points; \( P<0.001 \)) at the first assessment, and decreased to 2.11 points (95% CI=1.44-2.81 points; \( P<0.001 \)) at the end of the study | The effect of finasteride on sexual functioning is minimal for most men and should not impact the decision to prescribe or take finasteride | 1++               |

CI: Confidence interval, RCT: Randomized controlled trial

Use of finasteride and prostate cancer

In a double-blind, randomized multicentric prostate cancer prevention trial, finasteride 5 mg was compared with placebo to determine whether it reduced the risk of prostate cancer. The study reported that although finasteride prevents or delays the appearance of prostate cancer, it was associated with an increased risk of high-grade prostate cancer (evidence level 1+). Long-term follow-up of the same subjects showed that there was no significant difference in the rates of overall survival or survival after the diagnosis of prostate cancer (evidence level 1+). The increased incidence of high-grade prostate cancer in the finasteride group has been attributed to improved performance of prostate specific antigen screening.[29] Furthermore, studies

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Table 1: Studies which found the side effects to be not significant

| Study                  | Type of study/number of patients | Results                                                                 | Inference                                                                                              | Level of evidence |
|------------------------|---------------------------------|------------------------------------------------------------------------|--------------------------------------------------------------------------------------------------------|-------------------|
| Carbone et al.[17]     | Review of many RCTs - 73 papers | Problems of ejaculation (2.1-7.7%), erection (4.9-15.8%) and decreased libido (3.1-5.4%) | The population studied is different and hence generalization is not possible. So in spite of this being a review of RCTs, we grade it as low evidence | 1+                |
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| Moinpour et al.[21]    | Prostate cancer prevention trial. Randomized, double-blind, placebo-controlled, n=18,882 men | Finasteride marginally increased sexual dysfunction and its impact diminished over time; the increase in the sexual activity scale score relative to placebo was 3.21 points (95% CI=2.83-3.59 points; \( P<0.001 \)) at the first assessment, and decreased to 2.11 points (95% CI=1.44-2.81 points; \( P<0.001 \)) at the end of the study | The effect of finasteride on sexual functioning is minimal for most men and should not impact the decision to prescribe or take finasteride | 1++               |

CI: Confidence interval, RCT: Randomized controlled trial

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**Studies which found the side effects to be significant [Table 2]**

There are a few recent studies which have documented findings contrary to the above.[24-27] These suggest that

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Table 2: Studies which found the side effects to be significant

| Study                  | Type of study/number of patients | Results                                                                 | Inference                                                                                              | Level of evidence |
|------------------------|---------------------------------|------------------------------------------------------------------------|--------------------------------------------------------------------------------------------------------|-------------------|
| Carbone et al.[17]     | Review of many RCTs - 73 papers | Problems of ejaculation (2.1-7.7%), erection (4.9-15.8%) and decreased libido (3.1-5.4%) | The population studied is different and hence generalization is not possible. So in spite of this being a review of RCTs, we grade it as low evidence | 1+                |
| Mondaini et al.[18]    | 120 patients                    | The incidence of erectile dysfunction, decreased libido and ejaculation disorders were 9.6%, 7.7% and 5.7% for group 1 (5 mg finasteride without counseling) and 30.9%, 23.6% and 16.3% for group 2 (5 mg finasteride with counseling) | There was a significantly higher proportion of sexual dysfunction in patients informed about the sexual side effects (group 2) as compared to those who were not informed | 1+                |
| Kaufman et al.[19]     | 1553 patients                   | Decreased libido, erectile dysfunction and ejaculatory disorders occurred in less than 2% of men. The incidence of each side effect mentioned decreased to ≤0.3% by the 5th year of treatment with finasteride | Incidence of side effects was comparable to that of placebo, both at 1 year and at the end of 5 years | 1+                |
| Leyden et al.[20]      | Multicentric, double-blind, placebo-controlled study done at 45 locations followed by a 1-year open extension | Significant increase in hair count in the frontal scalp of finasteride-treated patients. Efficacy was maintained or improved throughout the 2nd year of the study. Sexual adverse effects were reported in approximately 2% of men in both treatment groups and they resolved while continuing in the study |                                                                                                          | 1+                |
| Moinpour et al.[21]    | Prostate cancer prevention trial. Randomized, double-blind, placebo-controlled, n=18,882 men | Finasteride marginally increased sexual dysfunction and its impact diminished over time; the increase in the sexual activity scale score relative to placebo was 3.21 points (95% CI=2.83-3.59 points; \( P<0.001 \)) at the first assessment, and decreased to 2.11 points (95% CI=1.44-2.81 points; \( P<0.001 \)) at the end of the study | The effect of finasteride on sexual functioning is minimal for most men and should not impact the decision to prescribe or take finasteride | 1++               |
associating prostate cancer with 1 mg of finasteride are lacking (grade of recommendation B).

**Position of other professional societies**
Given the conflicting data and importance of the subject, the International Society of Hair Restoration Surgery established a task force on finasteride adverse event controversies to evaluate published data and make recommendations.[28] The task force did not find much evidence on sexual dysfunction with once daily doses of 1 mg finasteride used for hair loss. They also urged the medical community to verify anecdotal reports and encouraged further studies.

**Food and Drug Administration position on the subject**
The Food and Drug Administration (FDA) recently reviewed 421 post marketing reports of sexual dysfunction with the use of finasteride, 1 mg submitted to the agency’s adverse events reporting system database between 1998 and 2011. They found that most of these side effects returned to normal within three months of discontinuing the drug.

In addition, analysis of controlled clinical trials showed that 36 (3.8%) of 945 men had reported one or more adverse sexual experiences during treatment with 1 mg finasteride as compared to 20 (2.1%) of 934 men who received placebo. The FDA reviewed 251 cases of altered semen quality with the use of 1 mg finasteride from the sponsor’s safety database and expressed the need for further evaluation of 13 cases.

**Food and Drug Administration recommendation on labeling changes**
On April 11, 2012, the U.S. FDA announced changes to the professional labels for Propecia (finasteride 1 mg) and Proscar (finasteride 5 mg) to include libido disorders, ejaculation disorders and orgasm disorders that continued after discontinuation of the drug. They also added a description of reports of male infertility and/or poor semen quality that normalized or improved after drug discontinuation. Although these are not established, they are included in labels to make patients and doctors aware of the side effect profile.

### Table 2: Studies which found the side effects to be significant

| Study       | Type of study/number of patients | Results                                                                 | Inference                                                                 | Level of evidence |
|-------------|---------------------------------|-------------------------------------------------------------------------|---------------------------------------------------------------------------|-------------------|
| Irwig et al.[24] | 71                              | 94% developed low libido, 92% developed erectile dysfunction, 92% developed decreased arousal and 69% developed problems with orgasm | New-onset persistent sexual dysfunction is significant and treating physician should discuss the potential risk of persistent sexual side effects. | 3                 |
| Mella et al.[8] | 12 studies                      | Increase in erectile dysfunction (RR, 2.22 (95% CI=1.03-4.78); P=1%; number needed to harm, 82.1 (95% CI=56-231)) and a possible increase in the risk of any sexual disturbances (RR, 1.39 (95% CI=0.99-1.95); P=0%). The risk of discontinuing treatment because of sexual adverse effects was similar to that of placebo (RR, 0.88 (95% CI=0.51-1.49); P=5%) (moderate equality evidence) | Oral finasteride increases hair count and increases the risk of sexual dysfunction | 1+                |
| Collodel et al.[25] | Report of 3 cases              | TEM analysis revealed altered sperm morphology consistent with necrosis and FISH data revealed elevated diploidy and sex chromosome disomy frequencies. This examination was repeated one year after the men had stopped using finasteride and were not receiving any other treatment. Recovery of the spermatogenetic process was observed. Motility and morphology improved whereas the meiotic pattern did not change presenting with elevated diploidy and sex chromosome disomy frequency | 3                 |
| Traish et al.[26] | Review                          | Prolonged adverse effects on sexual function such as erectile dysfunction and diminished libido were reported by a subset of men raising the possibility of a causal relationship | Discussion with patients on the potential sexual side effects of 5α-RIs before commencing therapy is ideal | 1+                |
| Gur et al.[27]      | Review                          | Prevalence rates of de novo erectile dysfunction are 5-9% | Study recommended that patients receiving therapy with 5α-RI should be counseled about the potential sexual and psychological adverse effects | 1+                |

TEM: Transmission electron microscope, FISH: Fluorescence in situ hybridization, CI: Confidence interval, 5α-RIs: 5 alpha reductase inhibitor, RR: Relative risk
Food and Drug Administration notification to healthcare professionals
The FDA states that there is no clear cause and effect relationship between finasteride and the sexual adverse events that continued after stopping the drug. Healthcare professionals should consider this new label information when deciding best treatment options. Finasteride remains a safe and effective drug for its approved indications.

Food and Drug Administration notification to patients
The FDA advised patients to consult their healthcare provider to discuss the risks and benefits of finasteride. They also urged patients not to discontinue the medication without first consulting the healthcare provider.

Position of IADVL therapeutic guidelines committee
In view of this, IADVL therapeutic guidelines committee makes the following recommendations to help members of IADVL in their practice and usage of the drug:

- Finasteride is effective in the management of androgenetic alopecia
- The mechanism of action and efficacy of this drug are proven
- Several studies have shown its safety over long-term use and no causal relationship has been confirmed between the drug and sexual side effects
- Researchers are yet to find safer and proven alternatives to finasteride
- Patients need proper counseling regarding the efficacy and side effects which needs to be provided in the form of patient information brochures
- The patients should contact the doctor for any advice, should he/she experience a side effect
- Most importantly, the intake of the drug is voluntary as patterned hair loss is only a cosmetic condition. It is entirely up to the patient whether to take the drug or not. If they choose to avoid the drug, they should be prepared for further progression of baldness
- The treating physician should provide full information about the drug in order to enable the patient to make an informed decision
- It is better to avoid the drug in patients who have had history of oligospermia or infertility, particularly if they are newly married and trying to raise a family. A patient who is anxious and expresses reservations about taking the drug also may avoid the drug. There is no recommendation that semen analysis should be carried out before prescribing the drug
  - No effort should be made by the physician to coerce the patient to take finasteride
  - The committee suggests that in patients who are apprehensive about the side effects, it is worthwhile to consider administration of lower or staggered doses of the drug to enhance patient compliance. The plasma half-life of finasteride is six to eight hours and the tissue binding is four to five days. A dose of 0.2 mg is adequate to suppress dihydrotestosterone levels in the scalp skin and serum. While daily doses of 0.2 mg caused 55% dihydrotestosterone suppression, 5 mg doses achieved 69% suppression. Efficacy has been demonstrated at all end points for finasteride at doses of 0.2 mg per day. Doses of 1 mg and 5 mg also demonstrate similar efficacy, but are superior to lower doses. The drug may therefore be initially administered at a dose of 0.5 mg daily for a short period. This would gain patient confidence and the 1 mg per day dose may be started once the patient is comfortable. However, the committee stresses that this is only an opinion of experts (grade of recommendation D).

Finasteride in the management of female pattern hair loss
Female pattern hair loss is difficult to manage and its exact etiopathogenesis is yet to be determined. Although finasteride has been shown to be effective in patients of female pattern hair loss, its clinical efficacy is controversial. There is limited data available on the subject, which is discussed below.

The mechanism of action of finasteride in female pattern hair loss is unclear. A daily regimen of 1 mg orally as indicated in male pattern hair loss may be recommended for those who fail or cannot tolerate minoxidil therapy. A trial of therapy for one year is needed to assess stabilization of hair loss and hair regrowth may take up to two years or longer. Although data is sparse, menopausal status, circulating androgen concentrations and concomitant symptoms of hyperandrogenism do not appear to predict a response to finasteride. The drug is generally well tolerated. Women of childbearing potential must adhere to reliable contraception while receiving...
finasteride. Pregnancy testing is mandatory before starting the drug. It is contraindicated in pregnancy due to teratogenicity.[32]

A study (evidence level 2+) which was double-blind and placebo-controlled evaluated the efficacy of finasteride in 137 postmenopausal women with androgenetic alopecia over a period of one year. They concluded there was no significant difference in the hair count between the finasteride and placebo groups at the end of one year. Both treatment groups had a significant decrease in hair count over the frontal and parietal scalp during the study period. The patient and investigator assessment, photographic assessment and scalp biopsy analysis did not demonstrate any slowing of hair thinning, increase in hair growth or improvement of hair appearance in the finasteride treated subjects when compared with the placebo group. In postmenopausal women with androgenetic alopecia, finasteride in doses of 1 mg per day taken for one year did not increase hair growth or slow the progression of hair thinning.[33]

Given the lack of efficacy of finasteride, 1 mg higher doses of 2.5 to 5 mg daily have been tried.[34,35] These studies concluded that oral finasteride in doses of 2.5 mg per day or more may be effective for the treatment of patterned hair loss in postmenopausal women, in the absence of clinical or laboratory signs of hyperandrogenism (grade of recommendation C).

**Therapeutic guidelines committee recommendations**

There is limited evidence (level C) for the use of finasteride at higher dosages for treatment of female pattern hair loss in postmenopausal women. The drug is a useful option for treatment failure cases. The use of this drug in childbearing age groups is not recommended in view of possible teratogenicity. It should be prescribed with proper counseling to avoid pregnancy. Informed consent and adequate contraceptive measures should be taken.

**Summary**

Finasteride is a widely used drug in dermatology for the treatment of androgenetic alopecia. Current evidence on the safety of finasteride indicates that it is safe, but there is growing concern about its sexual side effects. In view of this, proper information should be provided to patients before starting treatment.

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**Conflicts of interest**

There are no conflicts of interest.

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