The Floppy Iris Syndrome – What Urologists and Ophthalmologists Need to Know

Faruquz Zaman  Christian Bach  Islam Junaid  Athanasios G. Papatsoris
Jhumur Pati  Junaid Masood  Noor Buchholz
Department of Urology, Barts and the London NHS Trust, London, UK

Key Words
Floppy iris syndrome • Cataract complication • Tamsulosin • Benign prostatic hyperplasia • Alpha adrenergic blocker

Abstract
Introduction: Benign prostatic hyperplasia (BPH) and cataract formation are common in older people. Medical management of symptomatic BPH is often preferred to surgical treatment as surgery increases the risk of morbidities, whereas, surgery is the main form of treatment to restore sight in patient with cataract. The clinical treatment of BPH is either alpha-1 adrenergic antagonist alone or combination of alpha reductase inhibitor and alpha adrenergic receptor (AR) antagonist. There are four alpha-AR antagonists currently available to treat BPH. The uroselective alpha-blocker tamsulosin is the most commonly used drug among all. Studies showed that the majority of the patients who develop intraoperative floppy iris syndrome (IFIS) were on tamsulosin. Women are more likely to develop cataract than men and some recent studies showed that tamsulosin is effective in treating female lower urinary tract symptoms and thereby can cause IFIS during cataract surgery. Evidence Acquisition: We performed a critical review of the published articles and abstracts on association of IFIS with alpha-blockers and other medications as well as other medical conditions. Evidence Synthesis: Tamsulosin is the most common cause of formation of IFIS. However, not all patients given tamsulosin develop IFIS and cases have been reported without any tamsulosin treatment. Conclusion: Tamsulosin is a recognized cause to impede mydriasis and lead to IFIS during cataract surgery. Urologist should collaborate with their ophthalmology colleagues and general practitioner during prescribing tamsulosin in patients with history of cataract or waiting for planned cataract surgery. The increasing life expectancy and growth of older people will increase the number of men and women who suffer from lower urinary tract symptoms as well as cataract. Therefore, further research and studies are required to properly understand the relation of alpha blockers and IFIS.

Introduction
The intra-operative floppy iris syndrome (IFIS) is a variant of the small pupil syndrome. It has been observed during cataract surgery in some patients currently or previously treated with the α1 adrenoceptor (AR) antagonist tamsulosin. IFIS was first described by Chang et al. [1] in 2005. These patients tend to dilate poorly and to respond poorly to stretching during surgery. Without adequate pupil dilatation, IFIS may shrink the visualisation of the surgical field, including the cataract itself. This may impair removal of cataract and can lead to other complications such as rupture of the posterior capsule, which further increases the risk of other vision-threaten-
ing complications of cataract surgery [2]. Isolated cases of IFIS have been observed with other AR antagonists (alfuzosin and doxazosin), but to a lesser extent than for tamsulosin. However, reports have also been found in the absence of tamsulosin. IFIS is characterised by loss of muscle tone in the iris with symptom triad of pupil constriction despite pre-operative dilatation with standard mydriatic drugs, fluttering and bellowing of iris stroma, and a marked tendency for the iris to prolapse towards the side port incisions [1]. Ohtake et al. [3] showed that tamsulosin and other AR antagonists inhibit phenylephrine induced mydriasis and cause miosis to an equal extent and duration in an animal study.

**Discussion**

We know that age is a predominant risk factor for both benign prostatic hyperplasia (BPH) and cataract. Physicians can expect to see an increasing numbers of aging males on α1 AR antagonists who require operative intervention for cataract removal as α1 AR blocker is the most common drug for BPH. The other commonly used mediation for BPH is 5 alpha reductase inhibitors (5ARIs) which alleviates lower urinary tract symptoms by reducing the size of prostate over a period of a few months. There are two 5ARIs available for the treatment of BPH: finasteride and dutasteride. Finasteride inhibits only the type 2 isoenzyme of 5 alpha-reductase, whereas dutasteride blocks both, type 1 and type 2 isoenzyme of 5 alpha-reductase and thereby prevents conversion of testosterone to dihydrotestosterone more completely.

Patients taking α1 AR antagonists may be at risk of developing IFIS during cataract surgery. Recently urologists have begun to use tamsulosin to treat lower urinary tract symptoms in women as well. Therefore, ophthalmologists could expect to encounter this surgical problem more frequently with increasing use of this drug in the elderly female population that also require cataract removal. Moreover, roles have been suggested for the serotonergic and dopaminergic receptors and of the 1, subtype of the α-receptor.

In another prospective observational study conducted in a district general hospital in the UK, Amin K et al. [5] analysed a total of 1,462 cataract cases (1,267 patients) performed over 6 months. Of these, 23 eyes of 16 patients who were on tamsulosin were recruited into the study. All the patients were men with the mean age of 76 years. A small pupil was demonstrated intra-operatively in 69% (16/23) of the eyes. A floppy iris or iris prolapse during surgery was reported on 57% (13/23) of the eyes. Complications were posterior capsule rupture in 1 case (4%) and iris trauma in 5 cases (22%). The incidence of IFIS in patients undergoing cataract surgery over 6 months has been calculated therefore 0.9% in a representative UK patient population. It seems that IFIS have a strong association with tamsulosin use.

More recently, IFIS was reported to be 1.6% in a study population of 774 patients [6]. IFIS was documented in 14 of 18 patients (77.8%) taking tamsulosin. Consistent with the original report by Chang et al. [1], tamsulosin use overall was observed in 2.2% of patients in the age group prone for cataract.

Some studies have attempted to quantify the risk of IFIS with tamsulosin and to clarify the role of other α1 adrenergic receptor antagonists in IFIS. In a UK-based observational prospective study of 2,390 cataract procedures, Cheung et al. [7] identified 3 eyes with pure IFIS and 6 eyes with some features of IFIS out of 15 patients (17 eyes) given tamsulosin. Notably, the duration of tamsulosin did not correlate with the severity of IFIS.
Pathophysiology of IFIS
The most comprehensive review of adrenergic receptors in relation to the potential pathophysiology of IFIS was recently published [8]. Contraction of the iris dilator muscle via adrenergic stimulation results in mydriasis, which is necessary during cataract surgery. Therefore, agents such as topical phenylephrine, a α1 AR agonist, are routinely used in cataract surgery. Besides the effect of α1A AR on prostate tissue, several animal studies have isolated the α1A AR subtype as the mediator of iris smooth muscle dilatation [9]. It has been predicted that since tamsulosin is the only specific α1A AR antagonist marketed for BPH, it may also inhibit α1A receptor in the iris, thereby leading to IFIS. There are 3 subtypes of α1 ARs – α1A, α1B and α1D have all been cloned and characterised [10]. A fourth subtype α1L AR has been pharmacologically defined but not fully characterised or cloned. Tiwari et al. [11] have also proposed an important role for the α1L subtype receptor in mediation of iris dilatation in the human eye, and tamsulosin is considered to be a more potent antagonist of this receptor subtype than terazosin or doxazosin. As the exact mechanism by which tamsulosin induces IFIS has yet to be established, the role of other non-adrenergic receptors (i.e. dopaminergic and serotoninergic receptors) in inducing pupillary changes warrants consideration [12–16]. Tamsulosin seems to have a potent affinity for dopaminergic receptors as well [17].

Table 1 compares the mechanisms and effects of tamsulosin and other AR antagonists.

Evidence Acquisition
We performed a critical review of the published articles and abstracts on association of IFIS with alpha-blockers and other medications as well as other medical conditions.

Evidence Synthesis
Patients on Tamsulosin before Cataract Operation
In humans, tamsulosin is an irreversible antagonist of α1 adrenoceptors, and thus a withdrawal period of several days may not be sufficient to suppress the blockade of α1 adrenoceptors. There is little research between the length of time of tamsulosin use and the development of IFIS. Chang et al. [1] found that discontinuation of tamsulosin 4–7 days before surgery was helpful, but did not prevent IFIS completely. Settas et al. [21] reported no benefit from temporarily stopping alfuzosin treatment.

Association of the Floppy Iris Syndrome and Alpha Adrenergic Antagonists

Table 1. Comparison of effects of tamsulosin and other α-blockers

<table>
<thead>
<tr>
<th>Alpha-blockers</th>
<th>Receptor subtypes</th>
<th>Functions/Effects</th>
</tr>
</thead>
<tbody>
<tr>
<td>Tamsulosin</td>
<td>selective irreversible nor-adrenergic (NA) antagonist to iris dilator smooth muscle [18]. α1A + α1B, and α1L NA receptors [11] dopaminergic (D) receptors</td>
<td>potent affinity for D receptors greater affinity for α1A + α1B receptors than other α1 blockers α1L receptor might mediate iris dilatation tamsulosin is more potent antagonist of 1L receptor</td>
</tr>
<tr>
<td>Terazosin</td>
<td>non-selective competitive receptor subtype - α1A, α1B and α1D receptors</td>
<td>no selectivity towards α1A receptor which is thought to predominate in Iris α1A + α1B – uroselective α1B – vascular epithelium</td>
</tr>
<tr>
<td>Doxazosin</td>
<td>competitive non-selective antagonist</td>
<td></td>
</tr>
<tr>
<td>Prazosin</td>
<td>competitive antagonist in iris dilator smooth muscle [19]</td>
<td></td>
</tr>
<tr>
<td>Alfuzosin</td>
<td>competitive antagonist effect on iris dilator muscle has not been studied similar structure to prazosin and behaves as prazosin little/no affinity for S/D receptors [17] balanced binding affinity for the α1 receptor subtypes</td>
<td>similar uroselectivity to tamsulosin but behaves as competitive antagonist like terazosin and prazosin [20]</td>
</tr>
</tbody>
</table>
before surgery. Discontinuation of tamsulosin for 1 week or less before surgery might be insufficient because the elimination half-life for tamsulosin is about 48–72 hour. Chang et al. [1] also suggested that IFIS might be semi-permanent, and that blockade of α1A receptors in the iris dilator muscle could result in disuse atrophy of the muscle because IFIS was still evident in some patients despite drug withdrawal. Cheung et al. [7] postulates that if the proposed mechanism of disuse atrophy is correct one should expect a minimum time tamsulosin use before IFIS develops. In their prospective study looking at the prevalence of IFIS in patients taking tamsulosin, they noted that of the 17 eyes studied the shortest duration of tamsulosin use associated with IFIS was 3 months. However, Taben et al. [22] reported a case of IFIS after 1 day of tamsulosin (1 dose) therapy in patient with tamsulosin which may mean alpha 1 blockade alone without disuse atrophy could be enough to cause IFIS. Shah et al. [23] also published 1 case of IFIS occurring only 2 days after tamsulosin was started. Tamsulosin is often prescribed as a short term pharmacologic adjunct for the treatment of ureteric calculus. This is relevant because it appears that tamsulosin can cause IFIS almost immediately. In other patients however, pre-operative dilatation and iris flappiness seemed to improve after drug withdrawal. Chang et al. [1] recommended temporary dilatation and iris flappiness seemed to improve after drug withdrawal. Chang et al. [1] recommended temporary dilatation and iris flappiness seemed to improve after drug withdrawal. Michel et al. [24] have assessed the effects of α1 adrenergic receptor antagonists on pupil size and intraurethral pressure, and Schwinn et al. [25, 26] have depicted the effect of therapeutic blood level of any α1 adrenergic receptor antagonist given up to the day surgery. They advised that it would be prudent to stop all α1 adrenergic receptor antagonists before cataract surgery to avoid the possibility of IFIS, although the exact time and effectiveness of this action still need to be determined by careful recording of drug doses and plasma levels in future prospective studies.

Stoppage of tamsulosin 1–2 weeks prior to cataract surgery is anecdotally considered helpful, but the benefit and the duration of requirement of discontinuing the therapy prior to cataract surgery has not yet been established and remain unclear.

**Effects of Tamsulosin after Cataract Surgery**

There are only few studies which assessed the connection between tamsulosin exposure and post-operative complications. Scientists focused on 96,128 men aged 66 and older who underwent cataract surgery in Canada between 2002 and 2007. Of those, one group of 3,550 patients had taken tamsulosin while another 7,426 had taken different alpha blocking agent within 2 weeks of cataract surgery. In total 284 of the patients in the study experienced relevant side effects related to cataract surgery. The research found that patients who suffered complications were 2.3 times more likely to have taken tamsulosin than other alpha blockers, which showed no increased risk of eye injury. According to researchers [27], one in every 255 cataract surgery patients who take tamsulosin within 14 days of the surgery will develop severe eye complications and thereby increase the risk of needing second surgery. The authors also concluded that recent use of other alpha blockers and previous use of tamsulosin or other alpha-blockers had no significant effect on post-operative adverse events like retinal detachment, lost lens or lens fragment, or endophthalmitis.

On the other hand, Taben et al. [22] described in a case report that a patient had IFIS during cataract surgery 7 weeks after taking a single dose of tamsulosin. Eight months after tamsulosin use his pupils still dilated poorly, though slightly better than during surgery. This indicates that stopping the drug preoperatively may not decrease the incidence or severity of IFIS. Tamsulosin may bind α1 receptors on the iris longer than elsewhere. From an experimental study, it has been noted in Albino rabbits that IV alpha antagonists induced miosis was reversed 8 hours after cessation of the drug [24]. However, binding characteristics of alpha antagonists in human and darkly pigmented irises have not been well studied.

In contrast, Schwinn et al. [25, 26] considered the implications of withdrawing of α1-adrenergic receptor antagonist treatment and add that restarting of α1 adrenergic receptor antagonists immediately after cataract surgery seems carry little or no risk.

As the combination of cataract surgery and tamsulosin exposure is relatively common, patients should be properly informed of the risks of tamsulosin therapy, and during pre-operative assessments it is important to identify the use of tamsulosin by the patients as well as pre-operative assessment staffs.

**Comparison of Tamsulosin with other Alpha Blockers in Relation to IFIS**

Structural differences and well-defined mechanisms of action of AR antagonists might account for substantial differences in cases of IFIS. Alfuzosin has similar uroselectivity to tamsulosin and it inhibits completely and selectively α1 adrenergic receptors in the prostate, bladder base and prostatic urethra. Alfuzosin has been reported to show selectivity only for the α1A subtype [28] and act as a pure competi-
tive antagonist in human prostatic smooth muscle [29],
as do terazosin [30] and prazosin [20]. Similar to prazosin [19],
alfuzosin behaves as a competitive antagonist in human iris smooth muscle. Settas et al. [21] suggested
that the overall affinity of $\alpha_1$ adrenoceptor antagonists towards $\alpha_1$, receptors might explain IFIS.

Chang et al. [1] noted poor or moderately poor pre-operative dilatation, but no definite cases of IFIS, in the
group of patients taking prazosin, terazosin, and doxazosin. Evidence remains insufficient to suggest that IFIS is
always a side effect of $\alpha_1$ adrenoceptor antagonists. Isolated cases of IFIS associated with other $\alpha_1$ adrenergic
receptor antagonists have however been reported.

Another UK prospective study reported greater use of
doxazosin (n = 11) in 100 patients awaiting cataract surgery than of any other $\alpha_1$ adrenoceptor antagonists,
such as indoramin (n = 8), prazosin (n = 5), terazosin (n = 2), and tamsulosin (n = 3). Of this unselected popula-
nation, no patients developed IFIS and 1 had a constricted pupil [31].

A recent study directly compared the incidence of
IFIS attributable to tamsulosin with an active comparator
group [32]. In this retrospective study of 64 men totaling 92 eyes, there was an increased risk of IFIS in pa-

tients exposed to tamsulosin (86.4%) when compared to alfuzosin 15.4%. The adjusted odds ratio for IFIS in pa-

tients taking tamsulosin when compared to alfuzosin was
32.15% (95% CI). Furthermore, a fivefold increase in
surgical complication rates was observed in patients di-
agnosed with IFIS, highlighting its clinical significance.

Currently, the risk of IFIS has only been demonstrated
with systemic use of $\alpha_1$ AR antagonists. In a study com-
paring the incidence of IFIS between topical and system-
ic use of $\alpha_1$-AR antagonists, no cases were observed in patients taking bunazosin, a topical non-selective $\alpha_1$ AR
antagonist [33].

Relation of IFIS with $\alpha_1$, Antagonists and
Associated Diseases/Medications

Chadha et al. [34] assessed the analysed case of floppy
iris during cataract surgery and the use of $\alpha_1$ adrenocep-
tor antagonists and the presence of diabetes mellitus in
1,786 patients (1,842 eyes). Eleven eyes in 11 patients
had complete IFIS, and 18 eyes in 18 patients had in-
complete IFIS; 12 eyes in 21 patients given tamsulosin
had signs of complete or incomplete IFIS; however, 17
cases of IFIS (5 complete, none of whom had ever taken
an $\alpha_1$ adrenoceptor antagonists, and 12 incomplete – 1 of
whom was receiving doxazosin) were noted for patients
who were not given tamsulosin. None of the other pa-
tients taking doxazosin (n = 48), or those taking alfazosin
(n = 2) or terazosin (n = 1) had signs of IFIS. No relation
between diabetes and IFIS was found. The researchers
conclude that non-selective $\alpha_1$ adrenoceptor antagonists
are unlikely to be associated with IFIS, but they suggest
that other factors apart from tamsulosin may play an im-
portant part.

In a case report Taben et al. [22] described that a pa-

tient with a history of dermatomyositis developed IFIS
1 day after a single dose of tamsulosin. Similar findings
were noted intraoperatively 7 weeks later in the second
eye after no further tamsulosin therapy. A single dose
of tamsulosin and/or dermatomyositis may be associated
with IFIS.

In addition, Schwinn et al. [26] proposed that IFIS
may be associated with various diseases and medica-
tions. They mentioned that in animal studies, nitric oxide
has been shown to relax iris sphincter and dilator smooth
muscle. The patient who uses a nitro-glycerin patch daily
could be another potential cause of IFIS. However, there
are no clinical cases or human studies showing an asso-
ciation between nitrates and IFIS.

Furthermore, saw palmetto (Serona repens), a widely
used alternative therapy for BPH, was also associated
with IFIS in 2 patients [35]. Neither of these two patients
had taken prescription medications for BPH and they de-
velop moderate IFIS. Despite the development of IFIS,
the authors reported no significant surgical complica-
tions.

As we mentioned earlier, 5ARI, finasteride and dutas-
teride reduce prostate size by inhibiting conversion of
testosterone to dihydrotestosterone, taking 6 months for
a full therapeutic response. Two cases of IFIS were asso-
ciated with finasteride intake [36]. Neither of the patients
had taken systemic $\alpha_1$ AR antagonists prior to surgery,
and to date these are the only published cases associated
with 5ARI therapy.

There are some cases of IFIS have been linked to oth-
er medications including chlorpromazine, labetolol, and
donepezil, which have three distinct mechanisms of ac-

tion [37–39].

Conclusion

The $\alpha_1$ adrenoceptor antagonist tamsulosin is recog-
nised to impede mydriasis and increase the risk of IFIS.
However, the exact mechanism remains under investi-
gation. On one hand, not all patients given tamsulosin
develop IFIS; on the other hand cases have been reported

Association of the Floppy Iris Syndrome
and Alpha Adrenergic Antagonists

Curr Urol 2012;6:1–7

5
without any tamsulosin treatment prior to or after cataract surgery. Some patients with tamsulosin-related IFIS improve after drug withdrawal, whereas others keep having symptoms years after the treatment has stopped.

Patients should be educated and consented regarding potential risks of this drug class. The patient’s urologist should be prepared for possible modifications of their surgical technique. The urology community should be aware of the potential risks and should collaborate with their ophthalmology colleagues and general practitioners (GP) during prescribing tamsulosin in patients with history of cataract or waiting for planned cataract surgery. The increasing life expectancy and growth of older people will increase the number of men and women who would potentially suffer from LUTS as well as cataract. Therefore, further research and studies are required to properly understand the relation of alpha blockers and IFIS.

References


