Capsular block syndrome – report of two cases

Síndrome do bloqueio capsular tardio – relato de dois casos

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ABSTRACT

Two cases of late capsular block syndrome are reported. They were observed nine and five years after facoemulsification, in one eye with pseudoexfoliation syndrome and the other operated on for glaucoma with a functioning bleb. There was a large amount of milky material behind the intraocular lens, and reduction of the visual acuity (four lines of the Snellen chart in both cases). The condition was promptly resolved through minimal YAG laser posterior capsulotomy.

Keywords: Lens; capsule; Lens/surgery; Capsulorhexis/complication; Cataract extraction; Postoperative complications; Case reports

RESUMO

Relato de dois casos de síndrome do bloqueio capsular, de ocorrência tardia, nove e cinco anos após facoemulsificação, respectivamente, em um olho com pseudoexfoliação capsular e outro operado de glaucoma, com bolha funcionante. A condição se constituiu na retenção de grande quantidade de líquido esbranquiçado, atrás da lente intraocular, o que levou à redução da AV, de quatro linhas de Snellen, nos dois casos. Houve pronta resolução do quadro com mínima capsulotomia posterior.

Descritores: Cápsula do cristalino; Cristalino/cirurgia; Capsulorrexe; Extração da catarata/efeitos adversos; Complicações pós operatórias; Relato de casos

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INTRODUCTION

Capsular block syndrome occurs in the intra- or postoperative (PO) period; the PO form can be classified as early or late. Among the PO forms, the early type was the first to be described by Davison(1) in 1990. Some years later, in 1998, the late type was recognised as a variant of the same syndrome by Miyake et al.(2). It is also known as capsular bag hyperdistention, capsulorhexis block syndrome, capsular bag distension syndrome, or viscoelastic retention syndrome. Masket was the first to use the term capsular block syndrome (CBS) in 1993(3). It involves a continuous capsulorhexis and occurs due to adhesion of the anterior capsule (AC) to the intraocular lens (IOL)(4-6), preventing exchange between the inner capsular bag and the anterior chamber and leading to fluid retention behind the IOL. Although the process is generally related to IOL implantation in the capsular bag, it can also occur after IOL implantation in the sulcus. (6-9)

The early type is not rare. However, there are few reports of the late type. (1,5-8,10-16) Both types can impair visual acuity (VA) and lead to further complications that should be readily recognised and managed.

This paper reports two cases of CBS detected several years after cataract surgery, making considerations about its pathogenesis and treatment.

Description of cases

Case 1

The patient was a 78-year-old white male seen on January 4, 2005. He had undergone cataract surgery in the left eye (LE) five years earlier; there was no information about the surgical procedure or the type of IOL.

The patient has bilateral Fuchs dystrophy, with diffuse corneal oedema in the pseudophakic LE and compensated dystrophy in the phakic right eye (RE). The RE had cataract and pseudoexfoliation of the lens capsule. He underwent needle micropuncture in the LE, with significant relief of symptoms and improvement of VA to 20/70.

The remainder of the ophthalmic examination revealed no significant findings in both eyes (BE), and he was later submitted to triple surgery in the RE, without complications and with a good visual outcome. His condition remained unchanged until October 7, 2010.

On November 30, 2009 he returned complaining of worsening vision in the LE, starting less than two months earlier. His corrected VA was 20/150, but refraction could not be assessed because of media opacity. IOP was 14 mmHg. On examination under slight mydriasis (the pupil would not dilate much) the IOL was centred in the capsular bag and there was a large amount of white fluid occupying the space between the IOL and the posterior capsule (Figure 1A). The capsulorhexis rim was normal in the superior and temporal regions (the parts that could be observed with the mydriasis obtained). The anterior capsule was fibrotic in this area and was apparently closely adhered to the IOL. The anterior chamber was deep. There were also posterior subcapsular opacities similar to Elschnig spots (Figure 1B).

The patient received the diagnosis of CBS and underwent posterior capsulotomy. The fluid disappeared completely, rapidly dispersing through the anterior vitreous, with improvement of visual acuity to 20/70. The condition remained unchanged and this eye underwent corneal transplantation on May 14, 2010. On October 7, 2010 this eye had a corrected VA of 20/40 despite the graft sutures.

Case 2

The patient was a 76-year-old mixed-race female with nuclear cataract and glaucoma in both eyes; she was using travaprost. On March 26, 2004 she underwent combined phacoemulsification with IOL implantation and trabeculectomy in the LE, without complications. The IOL implanted in the capsular bag was an AR 40 from Advanced Medical Optics made of hydrophobic acrylic and PMMA haptics, with a biconvex 6-mm optic, a total diameter of 13 mm, and +22.00 D. The type of viscoelastic used in the procedure or the diameter of the capsulorhexis were not recorded.

In April 2006 the patient underwent needling with subconjunctival injection of mitomycin to recover a failed fistula, with successful restoration of drainage.

No abnormalities were noted until an evaluation on May 25, 2009, but in November 18, 2009, with a corrected VA of 20/30, an opacity was noted behind the IOL. This should have been clarified in an immediate follow-up visit for papillary dilation, but that was only done in June 2010, when the VA was already 20/50 with a refraction of -2.00 -1.75, 75°; the patient was using travaprost. On March 26, 2004 she underwent combined phacoemulsification with IOL implantation and trabeculectomy in the LE, without complications. The IOL had a -1.00 -1.50, 80° lens. The posterior capsule (PC) was distended, with an accumulation of white fluid between the PC and the IOL (Figure 2A); the anterior chamber depth was apparently normal for a pseudophakic eye.

The IOP was 11 mmHg, and there was a well-formed filtration bubble. A white mass was observed through the iridectomy in the superior periphery of the capsular bag (Figure 2B), as well as a posterior-inferior subcapsular spot-like opacity outside the dynamic pupillary area (Figure 2C). The edge of the anterior capsule had a significant degree of fibrosis and was apparently closely adhered to the IOL; the capsulorhexis diameter was 4.5 mm (Figure 2C).

Three days after YAG capsulotomy the corrected visual

Figure 1: A. Large amount of white fluid trapped behind the intraocular lens. The tortuous aspect of the anterior and posterior line of the fluid is due to the irregular corneal surface; thickened fibrotic anterior capsule; B. Posterior-inferior capsular opacity similar to Elschnig spots with direct oblique illumination.

Figure 2: A and B. Large amount of white fluid between the posterior capsule and anterior chamber behind the intraocular lens.
acuity was 20/20 with the old lens.

Similar to the LE, the RE had undergone combined surgery followed by needling, more or less at the same time, but had no particular changes; it also had a filtration bubble and controlled IOP without medication. However, in July 2007 it had undergone posterior capsulotomy.

**DISCUSSION**

Miyake et al.(5) proposed a classification for CBS in three types, depending on the time of onset: a) Intraoperative CBS: occurring in eyes with posterior capsule rupture and core dislocation blocking the anterior capsular opening after hydrodissection. It occurs more frequently in polar and mature cataracts and eyes with a greater visual axis(17,18) b) Early postoperative CBS: usually present on the first day(6,8,9,11,14,16) but may occur within the first two PO weeks. The seven cases described by Davison(11) were not observed on the first PO day, only when they returned for a follow-up visit two weeks later; c) Late postoperative CBS: tends to be a chronic process which manifests on average 3.8 years after surgery.

This classification, proposed by Miyake et al.(5) in 1998, was based on a review of 7 eyes with core dislocation after hydrodissection described by Hashimoto et al.(17); 13 early cases reported by Davison(11) and Holtz(10); and 44 late cases described by Ota et al.(18), Eifrig(19) and, most importantly, by Miyake et al.(5). Until a few years before that classification, only the early type had been recognised; then, in 1996, Robert Drews reported the intraoperative type in the Video Journal of Cataract and Refractive Surgery (cited in Miyake et al.(2)). The late type used to be considered as a different entity and was called secondary fluid cataract(5,18) or capsulorhexis-related lacteocrumenasia(19).

Many years later, in 2008, Kim et al.(13) proposed a new classification for PO CBS with three groups, taking into account its pathogenesis, based on the analysis of 8 cases: a) acellular, occurring early on the 1st PO day due to retention of viscoelastic, with little cell reaction or fibrosis between the IOL and the anterior capsule; b) inflammatory: cases of early CBS with inflammatory reaction and fibrin; c) fibrotic: cases of late CBS due to other mechanisms, such as metaplasia and proliferation of lens epithelial cells (LECs).

The early type seems to be much more frequent than the late type. In 10 articles published on the syndrome from 1992 to 2001 there were 41 reports of the early type(6,7,8,11,13,14) vs. 4 reports of the late type in 3 articles(8,12,14). However, the classification proposed by Miyake et al.(5) was based on a significant sample of 44 eyes with late CBS reported in only 3 papers.(5,18,19) Of these, 41 had been reported by Miyake et al.(5)

In early CBS, despite 6 reports of IOLs implanted in the capsular bag(13) and two reports of IOLs implanted in the sulcus(10) with an apparently normal anterior chamber, the anterior chamber depth is usually reduced.(1,2,5,11,14,16) In early CBS with IOL in the capsular bag, since there is no retraction of the capsular bag and because it has a much larger volume than the IOL, the lens tends to be pushed forward by the fluid that collects behind it, which can lead to myopisation as described by Davison(11) and Holtz(10). Therefore, any case of suspected myopic biometric error in the immediate PO period should be examined with pupil dilation to exclude CBS.(16) The same mechanism can also cause angle closure with increased IOP(12,5,10,14).

By contrast, in the late type this does not usually occur(2,5), but there tends to backward distension of the PC, as suggested by Theng et al.(14) and verified in the two cases reported here. Both eyes had a fibrotic ring around the capsulorhexis rim, and although there was a large accumulation of fluid behind the IOL, there was no significant IOL protrusion and the anterior chamber retained a depth compatible with pseudophakia. In only one case was it possible to assess refraction; a myopisation of 1 D was found, and as expected the IOP was normal. This generally occurs because in the late type capsular fibrosis tends to prevent the IOL from being pushed forward, probably not only due to the fibrotic ring which forms around the capsulorhexis rim but also to contraction of the capsular bag as a whole, with consequent tightening of the zonule.

This is generally expected in late CBS with IOL implantation in the capsular bag. However, in cases of IOL implantation in the sulcus, protrusion with myopisation and angle-closure glaucoma has been observed.(8) For the early type, Kim et al.(13) reported that increased axial length is a risk factor, mentioning a diameter greater than 25 mm. Although these cases have been reported with different lens types,(1,2,5,16) the early type is more common in lenses with flexible haptics, as they are more easily pushed forward.(5) In particular, the condition has been associated with a specific type of IOL, the Akreus Adapt by Bausch & Lomb. It has four points of support, a 6-mm optic made of hydrophilic acrylic, and no haptic angulation, which not only increases contact with the anterior capsule but also interferes with viscoelastic aspiration(16). Kim et al.(13) observed six cases in 206 cases (2.9%) with this IOL, while with other IOLs, according to various authors, the frequency ranges from 0.3 to 1.6%.(1,6,13,21,22) Marbeck et al.(16) reported an even lower prevalence of 0.02%.

In most reports of the early type the IOL was a single-piece Acrysof MA60BM made hydrophobic acrylic(5,6,11,13,14,15), but
according to Durack et al. CBS is independent of the type of material. However, such a relationship is unclear in the case of lenses with a biconvex design, which seems to facilitate closure between the capsulorhexis rim and the IOL. According to Davison(5) the Acriflavin IOL can cause CBS because of its shape, while PMMA IOLs can cause it because they produce a stronger reaction, with increased fibrosis and adhesion between the ante- rior capsule and the IOL. According to Holtz(10), CBS usually does not occur with lenses that have positioning holes, as well as smaller lenses and those with firmer angled haptics.

Other factors are the size of the capsulorhexis (small), the type of viscoelastic, and the amount retained in the eye.(2,4,6,13,15) All reports involved the use of sodium hyaluronate (Healon™ or Healon GV™), and in four cases an association of sodium hyaluronate with chondroitin sulphate (Viscoat®) was used.(3,14) Sodium hyaluronate retained in the capsular bag behind the IOL increases the osmotic gradient, attracting fluid that passes through the lens capsule, which acts as a semipermeable membrane.(10) This pushes the IOL forward toward the capsulorhexis rim, causing the blockade. This could not happen with viscoelastic of lower molecular weight, which would cause a lower oncotic pressure and would also be able to pass through the lens capsule, not being retained in the capsular bag.(10)

However, even in early cases, Davison(1) and Masket(4) minimise the role of the viscoelastic, emphasising the importance of retained lens materials such as subepithelial cells and their protein products, and possibly cortical remnants. This hypothesis is also cited in the early work of Miyake et al.(2,5)

Due to its pathogenesis, early CBS usually involves clear fluid without inflammatory reaction. However, Mardelli & Mehanna(20) described a case of early CBS with white fluid, classifying it as a sterile endophthalmitis due to cortex retention behind the IOL, but a with clean vitreous. This case could be included in the inflammatory group in the classification by Kim et al.(15), who reported two cases related to important anterior chamber inflammation and resolved with anti-inflammatory treatment.

As for late CBS, the accumulated fluid is white and milky, and Miyake et al.(5) suggest it is formed by LEC proliferation and metaplasia, with production of numerous types of collagen and extracellular matrix which accumulate in the capsular bag. Secondly, the process could be due to a difference in osmolality between the aqueous humour and the materials accumulated in the “closed chamber”, including residual LECs and their products and cortical remnants, which would attract aqueous humour.(10) Another possibility described by Nishi et al.(12) is that residual proliferating LECs become apoptotic and/or necrotic.

Eifrig(19) studied the fluid by electrophoresis and found a large amount of alpha crystallin and small amount of albumin, but no gamma globulin, which reinforces the hypothesis that the material is derived from the lens itself and is not due to an antigen-antibody reaction. A similar opaque fluid appears by liquefaction of the lens matrix in hypermature cataracts.(12)

LECs have a key role in the synthesis of lens protein matrix(23) and are found in the anterior capsule remnants in cataract surgery.(1) They must retain their synthesis ability, at least in part, in pseudophakic eyes, assuming that their products are constantly exchanged with the aqueous humour. Thus, according to Davison(1) and Holtz(10), AC adhesion with the IOL, which under normal circumstances occurs approximately two weeks after surgery, must not be complete, as if that happened the vast majority of cases would suffer capsular block with retention of fluid in the capsular bag. In both eyes of this report there were inferior subcapsular spot-like opacities, and in one of them a peripheral white mass could be seen through the superi- or iridectomy, resulting from LEC proliferation. These accumulations of proliferating cells could explain further production of the proteins mentioned above. However, in the large series of 41 eyes studied by Miyake et al.(5) capsular spot-like opacities were described in only 11 eyes, while 12 eyes had fibrotic opacities and one had a Soemmerring ring.

The two eyes reported here had IOLs in the capsular bag, which is the condition of most reported cases, both in early and late CBS.(1,2,6,11-15) However, there is one report of late CBS(8) and three reports of early CBS(6,7,9) with IOLs in the sulcus. Thus, regardless of the type of IOL implantation, the basic condition for CBS is occlusion of the anterior capsule opening preventing the exchange of fluid between the anterior chamber and the capsular bag.

In case 1 of this report, the phakic contralateral eye had capsular pseudexfoliation, a condition which has not been associated with CBS. In case 2, the eye with CBS had undergone combined surgery (cataract + IOL implantation + trabeculectomy), similar to the contralateral eye.

The interval between the procedures in both eyes was a few months. Both underwent needling to recover their fistulas, which had suffered failure. However, the eye without CBS had undergone posterior capsulotomy more than two years earlier. All procedures were uneventful. The association of CBS with glaucoma surgery was observed and supported by Muñoz-Negrete & Rebolloleda(13), who reported four cases (two trabeculectomies and two Ahmed valve implants). The authors suggest that these procedures, combined with cataract surgery, must be associated with CBS because the relative reduction of pressure in the anterior chamber can cause IOL displacement toward the anterior capsule. The cases were observed one month, five months, two years and three years after surgery. The authors point to the fact that the cases detected five months and two years after surgery probably had CBS since the immediate postoperative period, stating that the syndrome should always be kept in mind and should be included in the differential diagnosis of shallow anterior chamber after combined cataract and glaucoma surgery. Two other cases of cataract + glaucoma surgery were reported by Sorenson et al.(24) and McQuenn & Margo(25): one with trabeculectomy and the other with Ahmed valve implantation. In their large series with 41 eyes, Miyake et al.(5) make a correlation between CBS and glaucoma (four ca- ses), but also with other conditions that alter the blood-ocular barrier, such as diabetic retinopathy and uveitis (14 cases and 2 cases, respectively).

As to the time of onset of late CBS, it occurs on average 3.8 years(23) after cataract surgery, with reports ranging from 2 months to 6 years(3,5,8,12,14,15) According to Nishi et al.(12), fluid accumulation in late CBS slowly causes visual blurring after 5-7 years. Theng et al.(14) also believe that late CBS is subclinical in its early stages and is only detected later when PC opacification and impaired VA occur.

In this report, one case manifested more five years and the other nine years after cataract surgery. However, contrary to what is said by Theng et al.(14), in both cases VA reduction could not be explained by capsular opacity, because it was outside the visual axis. Therefore, the VA impairment of four lines in Snellen’s chart was related to the accumulation of large amounts of white fluid, which is consistent with the statement by Miyake et al.(5) that significant reductions in visual acuity related solely to fluid accumulation (which was observed in only 6 of their 41 cases) only occurs when the accumulation is large and the fluid is thick.
Regarding the therapeutic approach, in early CBS expectant management can be adopted at first, since spontaneous resolution is relatively common, being reported within 2 weeks to 2 months. In an early case followed by Durack et al. that received no treatment for 11 months, the patient remained stable throughout the period and was then treated by posterior capsulotomy. When spontaneous resolution does not occur, anterior capsulotomy can be used. When this is not effective or cannot be done because the pupil does not dilate, posterior capsulotomy is an alternative. Invasive treatments have been performed such as slit lamp needling and aspiration of material retained behind the IOL by irrigation/aspiration.

For late CBS, there are only 3 reports of spontaneous resolution. The generally indicated procedure is posterior capsulotomy because it promotes greater IOL stability in the capsular bag and has a lower risk of complications. There is only one report of a case that did not resolve with the posterior YAG capsulotomy, where the patient was submitted to surgical capsulectomy with posterior vitrectomy.

In both cases reported here, a minimal opening was done in the PC, because there was no capsular opacity corresponding to the dynamic pupil area. It was also carried out with a very low power, aiming only to release the fluid and trying to prevent this initial opening from pushing the PC due to the unknown pressure of the large amount of fluid, causing a large rupture and jeopardising IOL support and maintenance of the vitreous in its place. Contrary to what was pointed out by Muñoz-Negrete & Rebolloda, despite the large amount of white fluid trapped behind the IOL we had no difficulty reaching the PC, which was achieved with a single laser shot. Also, as expected, there was no inflammatory reaction in the eye as a result of fluid extravasation into the anterior vitreous.

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