Surgical management of anomalous origin of coronary artery from pulmonary artery

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Background
Anomalous origin of coronary artery from pulmonary artery (AOCAPA), as is evident from the name, is defined as abnormal origin of either coronary artery from the pulmonary artery. The consequences vary in most cases and these anomalies lead to severe coronary hypo-perfusion and ventricular dysfunction. The common variants of this cardiac malformation are an anomalous origin of a left coronary artery from a pulmonary artery (ALCAPA) and anomalous origin of the right coronary artery from a pulmonary artery (ARCAPA). Another rare variant is left main coronary artery atresia that resembles ALCAPA in its mode of presentation. This article presents a single surgeon experience of managing this complex subset of the coronary anomaly from April 2006 to July 2019.

Material and methods
The 105 patients, who underwent surgery for AOCAPA from April 2006 to July 2019, have been included in the study. The patients have been analysed by follow-up echocardiography and electrocardiography (ECG) at our hospital by paediatric cardiologists. Out of 105 patients of AOCAPA, 98 (93.3%) patients underwent ALCAPA repair, of which 59 (60.2%) were males and 39 (39.7%) were females. Four out of five patients, who had an anomalous origin of the left coronary artery from the right pulmonary artery (ALCARPA), had an intramural aortic course. Three patients (3%) had left main coronary artery atresia and four patients (4%) had ARCAPA. It may be mentioned that seven infants (7.14%) and one adult patient (1%) underwent concomitant mitral valve repair. All the patients with ALCAPA, left main coronary artery atresia and ARCAPA, and 1 of the patients with ALCARPA, underwent coronary relocation. In four out of five patients with ALCARPA, unroofing of intramural course was performed.

Results
Out of 105 patients of AOCAPA, 9 (8.5%) patients had in-hospital mortality. Five infants (5.0%) with ALCAPA and one patient (1%) with ALCARPA died in the post-operative period due to severe left ventricular dysfunction, mitral regurgitation (MR) and sepsis. One adult patient (1%) with ALCAPA, who underwent coronary relocation using in situ trap door technique and mitral valve (MV) repair, died due to massive intracranial bleeding. Two patients out of three (66.6%) with left main coronary artery atresia died in intensive care unit (ICU) after 3rd and 4th postoperative day, due to low cardiac output, severe ventricular dysfunction and severe MR. Patients were followed up for a median 5.9 years. Seven patients were lost to follow-up, including the sole survivor of left main coronary atresia, after a median follow-up of 4 years after surgery. Three patients underwent mitral valve replacement for progressive residual MR. There has been no late mortality.

Conclusion
AOCAPA is a rare congenital cardiac anomaly, which usually presents in infancy with left ventricular dysfunction and mitral valve regurgitation. Early diagnosis and surgical re-establishment of the dual coronary system has given gratifying results, with improvement in left ventricular function in survivors. Mitral valve intervention for MR was required, in both early and late phases.

Keywords
AOCAPA: Anomalous origin of coronary artery from pulmonary artery · ALCAPA: Anomalous origin of left coronary artery from pulmonary artery · ALCARPA: Anomalous origin of left coronary artery from right pulmonary artery · ARCAPA: Anomalous origin of right coronary artery from pulmonary artery · Left main atresia · Congenital coronary anomaly

Introduction
Anomalous origin of the coronary artery from the pulmonary artery (AOCAPA) is a rare congenital cardiac anomaly, with an incidence of 0.01% [1] in the general population. Anomalous
origin of the left coronary artery from the pulmonary artery (ALCAPA) is the commonest, accounting for 1 in 3,00,000 live births [2]. It can easily be diagnosed in infancy, as the patient shows clear-cut symptoms. The problem calls for timely treatment, as almost 90% of patients succumb to the ailment in the first year itself [2, 3]. However, a fraction of patients with ALCAPA may reach the late adult stage, if either the symptoms are dormant, or they may present with symptomatic myocardial infarction (MI) or as severe mitral regurgitation (MR), due to chronic ischemia. Anomalous origin of the left coronary artery from the junction of the main and the right pulmonary artery (ALCARPA) is also a rare form of ALCAPA. In this, the left coronary artery arises from the junction of the main and right pulmonary artery (PA) and enters into the aorta with an intramural or extramural (Fig. 1) course in the aortic wall. It exits through the normal course of the left main coronary artery [4]. Left main coronary artery atresia is also an uncommon anomaly, in which there is no left coronary ostia and the proximal left main coronary trunk ends blindly. The blood flows from the right coronary artery (RCA) to the left, through small collateral arteries in retrograde manner, in at least one of the left side arteries. Its clinical presentation is similar to ALCAPA with extensive collaterals in the right ventricle and area around the main PA [5]. Anomalous origin of the RCA from the pulmonary artery (ARCAPA) accounts for 0.002% of cases with congenital heart disease (CHD) [6]. It is a relatively benign condition and presents usually in adolescence and adults. Almost 60% of ARCAPA patients are asymptomatic and have the best prognosis amongst all forms of AOCAPA [6, 7].

**Material and methods**

A retrospective case records review was made of all patients with diagnosis of AOCAPA. During the study period, April 2006 to July 2019, 105 patients underwent surgery for AOCAPA, 98 patients (93%) had ALCAPA, including 5 patients (4.7%) with ALCARPA; 3 patients (3%) had left main coronary artery atresia and 4 patients (4%) had ARCAPA. The median age at presentation was 5.8 months (range 14 days to 45 years). There were two adult patients (2%). Fifty-nine were males (60%) and 39 (40%) were females. Three (3%) had left coronary artery (LCA) arising from non-facing sinus (Fig. 2). Twenty-five (25.5%) of the 33 (33.6%) patients underwent surgery in early infancy (<6 months), and 51 (52%) patients in late infancy (>6 months). Seven patients (6.8%) out of 98 with ALCAPA had associated lesions. Two patients (2%) had secundum atrial septal defect (ASD), while one (1%) had primum ASD with a double orifice mitral valve (MV). This patient underwent MV repair and closure of the ostium primum ASD. One patient (1%) had large patent ductus arteriosus (PDA) with severe pulmonary arterial hypertension (PAH), and one 3-month-old child had a large aorta-pulmonary (AP) window with left coronary artery arising from the PA side of the window (Fig. 3). He had good left ventricular (LV) function with mild MR. One patient (1%) had a large ventricular septal defect (VSD) with severe PAH. One (1%) of the patients with ALCAPA had accessory left anterior descending (LAD) coronary artery from RCA (Fig. 4), and he had mild MR and moderate LV dysfunction. Seven infants (6.6%) and two (2%) adults ALCAPA patients had severe MR needing MV repair or replacement. Three patients (2.94%) had left main coronary artery atresia and 4 patients (3.8%) had ARCAPA (one of them had an intramural course in PA with moderate size secundum ASD).

12-lead electrocardiography (ECG) and transthoracic echocardiographic examination was performed on all patients. Ejection fraction (EF) and fractional shorting (FS) were employed to classify LV systolic dysfunction. According to this, LV systolic function was accepted as normal when LVEF was ≥60%, and there was mild, moderate and severe systolic dysfunction if LVEF was 50–59%, 35–49% and less than 35%, respectively. Mitral regurgitation was graded as I (none/trivial), II (mild), III (moderate) and IV (severe). Coronary angiography and computed tomography (CT) pulmonary angiogram were performed in selected patients. All patients with ALCAPA, ARCAPA, and 2 patients with left main coronary artery atresia underwent left main coronary artery transfer to the aorta. Four patients (3.8%) with ALCAPA underwent unroofing of an intramural intra-aortic course and one patient with extramural course underwent coronary relocation (Fig. 1). None of the ALCAPA patients were put on extracorporeal membrane oxygenation (ECMO) support in the immediate post-operative period. The data on follow-up, with last clinical, ECG and echocardiography evaluation, and any further surgical intervention, were noted.
Diagnosis was the indication for surgery. Patients in sepsis were stabilised preoperatively. Chest was opened through midline sternotomy. Pericardium was opened. Collaterals between aorta and PA groove were cauterised. Branch PAs were dissected and looped, PDA was dissected. Using standard technique of caval cannulation, PDA/ligamentum arteriosus was ligated and divided. Cardioplegic arrest was achieved using ante-grade cardioplegia via aortic root, after occlusion of branch PAs. Left atrial (LA) vent was placed through patent foramen ovale (PFO). Main PA was incised just below the level of right PA. After division of anterior half of the main PA, position of the coronary artery ostium was confirmed (just to avoid any injury to coronary artery in case of misdiagnosed ALCAPA). Main PA was transected, position of the anomalous artery was confirmed and coronary button was excised for transfer to aorta (Fig. 5a). Coronary button was dissected off the epicardium for a centimetre to avoid any unnecessary traction on the coronary button artery. Collateral branches, which were divided during dissection of the coronary button, were cauterised or sutured with 7-0 polypropylene. Defect in the main PA was repaired with autologous glutaraldehyde-treated pericardium (Fig. 5b). A stay stitch (Fig. 5b) was taken at the expected site of transfer, at postero-lateral aspect of aorta at nearly 7 o’clock position, which is usually slightly difficult for the surgeon to see. Stay suture was pulled towards the first assistant and the aorta was cut opened at the site of stay suture.
The opening was inspected and was further enlarged using a punch. The coronary artery was transferred using 7-0 polypropylene continuous suture (Fig. 5d). We never had any injury to aortic valve leaflet or the commissure. Although we have only two adult ALCAPA in our series, we believe that the direct transfer of coronary artery is possible in adult patients using in situ lateral or medial base trap door technique or by using native tissue for coronary artery reconstruction [8]. Small dose of cardioplegia was given to check for any bleeding from anastomotic site or from any collateral vessel. We usually do MV repair only if there is some organic cause for MR, like cleft or prolapse, or in cases of moderate to severe MR, using posterior suture annuloplasty. The other associated lesions are managed after coronary transfer (MV repair, VSD, ASD closure). In one of our patients, who had a large AP window (Fig. 3), the patch between the aorta and PA was placed in such a way, as to leave the left coronary artery towards aortic side of the patch. Pulmonary artery posterior layer was constructed on clamp and remaining anterior part was reconstructed off-clamp (Fig. 5e). The bypass was weaned off using adrenaline, milrinone or levosimendan support. We usually do not place LA line as a routine and we measured LA pressures and performed transesophageal echocardiography (TEE) only when either MV repair was performed, or if there was difficulty in weaning from bypass. It was not uncommon to have borderline haemodynamics after coming off bypass which gradually improved with cardio pulmonary bypass (CPB) support. Occasionally, there was some bleeding from left postero-lateral aspect of main PA, which usually settled with protamine administration or glue application. We preferred to close the chest after surgery, and chest was only left open, to be closed later, if there was medical bleeding or borderline haemodynamics.

**Surgical technique for ALCARPA (Fig. 6a–e)**

If the diagnosis is known before surgery, the entry of the left coronary artery, from the junction of main PA or right PA to
posterior surface of ascending aorta, can be seen (Fig. 6a, b, d). After giving cardioplegia from ascending aorta and main PA, after occlusion of branch PAs, the aorta was transected just at the level of right PA. It is difficult to understand the intramural intra-aortic course of left main coronary artery from inside the aorta, so it is safer to open the right PA and identify the opening of coronary artery from right PA and the entry point in aorta. The coronary artery can be disconnected from right PA and its opening can be unroofed using Potts scissors (Fig. 6c), until the lower end of its intramural course, which is usually a bigger opening of the left main coronary artery. The unroofed edges are re-endothelialised (Fig. 6e). The junction of main PA and right PA can be reconstructed using pericardial patch and the aorta can be reconstructed. If the course of ALCARPA is extramural, then it has to be relocated.

Surgical technique for left main coronary artery atresia is demonstrated in Fig. 7a–e. Left main coronary atresia should always be suspected first, if the anomalous LCA is not seen to be arising from the main PA during surgery and the left main coronary artery should be looked for behind the main PA. In our experience, it usually ran behind the main PA and ended blindly as a bulbous end (Fig. 7a), surrounded by extensive collaterals, just 3–4 mm short of postero-lateral aspect of the aorta from where the left main coronary artery arises (Fig. 7b). If the diagnosis was known before surgery, then after giving cardioplegia through aortic root, the main PA was dissected, as is done in ALCAPA. The bulbous portion of the left main coronary artery was identified and dissected off the epicardium for about a centimetre, and the blind end was cut opened (Fig. 7c, d). Usually, the cut end was slightly thickened and could easily be sutured to the aorta at 7 o’clock position, like proximal coronary anastomosis (Fig. 7e). Small dose of cardioplegia is given through the aortic root and flows through the coronary artery and any bleeding from collaterals was checked and fixed at the same time. Usually, there were a lot of collaterals around the left main coronary artery and special attention was required to control bleeding from collaterals. In our experience, patients with left main coronary atresia were
associated with severe MR and we did posterior suture annuloplasty in both our patients. Main PA can be reconstructed as described in ALCAPA (Fig. 7e). These patients may need prolonged bypass support or may even need ECMO support (we did not have access to ECMO, when we operated on these patients).

Surgical technique for ARCAPA (Fig. 8a–c)

Standard CPB and cardioplegic arrest, by giving cardioplegia through both aorta and main PA (well above the origin of RCA), after occlusion of branch PAs, were utilised. RCA button was excised and the resulting defect was repaired using a glutaraldehyde-treated pericardial patch (Fig. 8a, b). The coronary artery was mobilised about a centimetre and was transferred using punch hole technique to ascending aorta (Fig. 8c). The associated lesions were treated at the same time and bypass was weaned off in a routine fashion. As the myocardium is well preserved, these patients behave very well in the post-operative period and can be weaned off ventilator within a few hours.

Statistical analysis

All the statistical analyses for the study were performed using SPSS software v 20.0 (Chicago, IL, USA). The continuous and normally distributed data was expressed as mean ± SD, whereas categorical data was presented as frequency and percentage. As age of the patients at the time of operation and follow-up duration of the patient population were heterogeneous and non-normally distributed, they were expressed as median and range. Survival analysis was also undertaken and is presented as Kaplan-Meier survival curve. The patients were grouped according to the age at the time of operation, and the improvement in LV function and MR during post-operative follow-up, according to the groups, was assessed and plotted.
Results

Patient characteristics

Out of 105 patients of AOCAPA, 98 (93.3%) patients underwent ALCAPA repair. Median age at operation was 5.8 months. Four (4%) out of five patients had ALCARPA with intramural aortic course. Three patients (3.06%) had left main coronary artery atresia and four (4%) patients had ARCAPA. Seven infants (7.14%) and one (1%) adult patient underwent concomitant MV repair. One adult patient underwent MV replacement.

The patients, presenting in early infancy (<6 months), had mean LVEF of 43.5% ± 5.1% and those presented in late infancy (>6 months) had mean LVEF of 38.6% ± 4.6%. Twenty-five (25.5%) of the 33 (33.6%) patients underwent surgery in early infancy (<6 months), 25 (25.5%) had grade II to III MR and 8 (8.1%) patients had grade III to IV MR. Of the 51 (52%) patients who underwent surgery in late infancy (>6 months), 38 (38.7%) had grade III to IV MR and 13 (13.2%) patients had grade IV MR.

Mean CPB and aortic cross-clamp time were 93 ± 9 min (81–102 min), and 63 ± 7 min (55–72 min) respectively.

On follow-up, all patients underwent ECG and echocardiographic examination.

Early mortality (Fig. 9)

Early mortality was 8.5% (9 out of 105) in the entire cohort. Five (4.7%) infants with ALCAPA and one (0.95%) patient with ALCARPA died in post-operative period due to severe LV dysfunction, MR and arrhythmias. One (0.95%) adult patient with ALCAPA died due to massive intracranial bleeding. Two (1.9%) patients with left main coronary artery atresia died in intensive care unit (ICU), after 3rd and 4th post-

Fig. 8 Schematic diagrams showing surgical technique for ARCAPA. a Showing ARCAPA. b Showing RCA button harvesting. c Showing relocation of RCA and pericardial patch reconstruction of MPA
operative day, due to low cardiac output, severe ventricular dysfunction and severe MR.

**LV function (Fig. 10)**

Median follow-up of ALCAPA patients was 5.9 years. One newborn, who had moderate LV dysfunction and mild MR, had normal LV function and no MR after 6 months of surgery. Thirty-three (33.6%) patients, who underwent surgery in early infancy (<6 months), had good recovery of LV function (mean LVEF 50 to 55%) and fractional shortening of 25 to 28%. Fifty-one (52%) patients, who underwent surgery in late infancy, had impairment of LV function (mean LVEF 40 to 50% and fractional shortening of 20 to 25) and 14 (14.2%) operated after infancy, without any left to right shunt but with severe ventricular dysfunction, had persistent mild to moderate LV dysfunction with gradual improvement of LV function (mean LVEF between 40 and 45% and fractional shortening 18 to 21%).

**Mitral regurgitation (Fig. 11)**

Thirty-three (33.6%) patients, who underwent surgery in early infancy with moderate MR, continued to have mild regurgitation on follow-up. Fifty-one (52%) patients operated after 6 months of age continue to have mild to moderate MR on follow-up. Fourteen (14.2%) patients, who were operated after infancy, continue to have mild to moderate MR on follow-up and three of them underwent MV replacement after 3 years, for unrepaired progressive severe MR.

We have started repairing MVs in patients with severe MR presenting in late infancy, after July 2012. Seven patients (6.86%) presented after infancy. They had severe MR before surgery and had undergone MV repair, but continued to have mild to moderate MR and were on medications.

**Presence of left to right shunt**

Six patients (5.7%) with left to right shunt and one (0.9%) with accessory LAD from RCA had normal preoperative LV function and no MR. This was no doubt secondary to severe pulmonary hypertension limiting the retrograde steal from the anomalous LCA.

**Follow-up**

Eighty-five patients were followed up until July 2019. Median follow-up was 5.9 years (1.5–12.5 years). Seven patients were lost to follow-up after a median follow-up of 4 years (3.2–6 years). Three patients underwent MV replacement for severe progressive unrepaired MR. All patients were in sinus rhythm, and no ischemia or arrhythmias were identified. None of our patients had aortic coronary stenosis. No late deaths were reported in any patient with AOCAPA, to date.

**Discussion**

Anomalous origin of the coronary artery from the PA is a rare congenital cardiac anomaly, with an incidence of 0.01% [1] in
the general population. ALCAPA is the commonest, accounting for 1 in 3,00,000 live births. It can easily be diagnosed in infancy as the patient shows clear-cut symptoms [2]. The problem calls for timely treatment, as almost 90% of patients succumb to the ailment in the first year itself [2, 3].

ALCAPA patients are usually infants, who after birth register decreased pulmonary vascular resistance causing left to right shunt [2]. We have operated only one 14-day-old baby, who had mild preoperative MR and moderate LV dysfunction with EF of 45%. One of our patients, a 3-month-old infant, was diagnosed to have ALCAPA during surgery for the AP window. The patch was placed in such a manner that the LCA was placed on the aortic side of the patch. Only 11 cases of ALCAPA with AP window have been reported to date in English medical literature [9]. Several techniques for surgical management of ALCAPA have been reported since Sabiston and associates described ligation of the LCA at its origin [10]. Neches and colleagues [11] first time reported direct implantation of left coronary into the aorta, with a good result, and it is now universally accepted. We also believe that direct aortic implantation is the treatment of choice for ALCAPA in infants [12], even if the coronary artery is arising from non-facing sinus. Preservation of two coronary system helps in early recovery of LV function. In infants, it is possible to mobilise the

**Fig. 10** Time course for left ventricular ejection fraction change in ALCAPA patients

**Fig. 11** Time course for MR in ALCAPA patients
coronary and directly anastomose it to the aorta, on its postero-lateral aspect. Direct aortic implantation has been shown to have a higher rate of late patency [13]; Ben Ali et al. had demonstrated that 2 of their 61 patients required re-operation for coronary stenosis after direct aortic implantation [14]. Post-operative coronary stenosis should be suspected in a patient with persistent LV dysfunction or worsening MR, although we have not come across any such incidence in our series.

Our experience with ALCAPA in adult patients is limited, as most of these are managed by cardiac surgeons for adults, and they usually perform coronary artery bypass surgery (CABG) with ligation of proximal left circumflex (LCX) coronary artery for such patients [15, 16]. In our two adult patients, we could mobilise the coronary and could relocate it on aorta, using the trap door technique. However, studies suggest that Takeuchi technique is a better technique for ALCAPA in adult patients, especially when the left coronary artery is arising from the non-facing sinus and in adult patients where mobilisation is difficult [17]. However, late baffle leak and pulmonary stenosis each have high incidence, necessitating lifelong care. Even baffle obstruction and aortic regurgitation have been reported [17]. We have no practical experience with Takeuchi’s operation.

**Left ventricular function**

LV dysfunction has been reported as the main risk factor for perioperative mortality [18, 19]. We have observed that there is the minimal improvement in LVEF on the day of surgery. However, in most of the patients operated in early infancy, improvement in LV function and ejection fraction was seen in a week or so. In the patients operated in late infancy, LV function may not improve at all, even at the time of discharge. Improvement in LV function is ultra-slow in the follow-up period of the first 3 months. Patients operated in early infancy or as newborns have better post-operative recovery and speedy recovery of their LV function on follow-up [20, 21]. Despite good clinical condition and normalised LVEF (more than 55%) in ALCAPA patients after surgical repair in the long-term follow-up, the diastolic and longitudinal systolic function and LV torsion of all cardiac chambers remained impaired, especially in the LCA region. Lifelong surveillance is mandatory [22, 23], as ischemia can lead to myocardial scarring, even in early infancy. The recovery of LV function is closely related to scar burden [24]. In our experience, except for the patient who had left to right shunt, severe PAH and accessory LAD, patients operated within 1 month of birth or early infancy had the best outcomes. In contrast to the series reported by Monge et al. [25], we never came across any patient who underwent surgery in late infancy and had normal LV function during follow-up. Probably this is because of late diagnosis and delayed presentation in developing countries, like ours [18]. However, most of our patients showed gradual improvement in LV function and symptomatic improvement on after-load reducing medicines.

**Mitral regurgitation**

MR is an important part of the disease and, to an extent, denotes the severity of ischemic damage of the LV. However, controversy still exists over the initial management of MR, with some groups recommending MV repair at the time of ALCAPA repair [24, 25]. Various studies have suggested that the cause of MR is LV ischemia and in the majority of patients, MR improves after the surgery.

Most centres do not recommend MV repair during initial operation, owing to the fear of prolongation of the cross-clamp time in the setting of an already-compromised myocardium. Medical practitioners generally believe that with the improvement of LV function and dimensions after successful revascularisation, the mitral annular size shall decrease with subsequent improvement in MR. However, our experience does not corroborate this view point, and for patients who come in late infancy and childhood with severe MR, where the cause of MR is ischemic lesions of the papillary muscle (organic MR), we recommend MV repair [18, 19]. In this group, mitral annuloplasty in association with coronary reimplantation is indicated due to the reduced likelihood of regression of MR after coronary revascularisation [26]. If the MV is not repaired in these patients during surgery, their post-operative course is difficult and most likely will need reoperation for MR. Severe MR leads to prolonged ventilation and it is extremely tough weaning from the ventilator. So, since 2012, we electively repair MV in patients with moderate to severe MR using posterior suture annuloplasty [18, 20, 26]. We have observed that in the patients, who underwent MV repair for severe MR, even if there was residual mild to moderate MR, their post-operative course in ICU was smooth. Although most of the patients regain their LV function and there is improvement in the degree of MR with coronary revascularisation, few patients still show the sequel of myocardial damage and a hemodynamically significant degree of MR, requiring close follow-up and further surgical intervention. Reduction annuloplasty, with or without chordal shortening and quadrangular resection, is the procedure of choice for the ischemic MR in children. Otherwise, MV replacement with mechanical prosthesis should be the last resort for patients with MR refractory to conventional repair [21].

For ALCAPA, we have reviewed and summarised cases reported in the literature [4]. Coronary reimplantation without unroofing and unroofing alone have their advantages and disadvantages. We found that the unroofing technique, which we have used in our patients, is simple and reproducible. When encountered, unroofing is always a preferred technique, as compared to reimplantation. The main problem of reimplantation
technique for ALCAPA is the risk of “kinks” of the coronary artery as the left coronary button has to be rotated almost 90–180 degrees for reimplantation. The transfer site would be far above the sinus of Valsalva, almost at the mid-section of the ascending aorta, which adversely affects LCA blood pattern, as compared to the case whose LCA has the origin inside the sinus [27]. The consistent presence of an intramural segment in the intramural coronary artery is known to pose potential risks of myocardial ischemia, and even sudden death [28]. The unroofing procedure would seem to be superior to reimplantation, because it can normalise virtually the entire course of the LCA, without leaving the intramural segment, which could eliminate a potential nidus for lethal events and it is technically simple too. We perform unroofing with the same technique that we had described for transposition of great arteries with intramural coronary artery [29]. We believe that unroofing leads to uniform coronary flow from the sinus of Valsalva.

Imaging almost always fails to diagnose intramural course. Echocardiogram fails to detect the aortic intramural course, due to a vertical course of the anomalous LCA parallel to ascending aortic wall, with the intimal relationship between the aorta and anomalous LCA. CT angiography also failed to diagnose the intramural aortic course in two of our cases. However, on reviewing the literature, indicators of latter are as follows: (1) high origin of the LCA at the pulmonary trunk or on the right PA, (2) ascending and parallel course of the ALCAPA to the aortic wall and (3) presence of a left perpendicular angle formed by the junction of the vertical intramural aortic course [30, 31]. It is not that ALCAPA always has an intramural course. Rarely, the LCA may arise from right PA with the direct extravascular course and in such a situation, one has to relocate the coronary artery by turning the ostium by 180 degrees. Even single coronary artery origin from right PA, without any intramural course, has been reported in English medical literature [32, 33].

Both infants and adult patients with left main coronary artery atresia have been reported [5, 34, 35]. Adults may present as patients of ischemic heart disease and only during angiography may be diagnosed to have left main atresia. A 76-year-old asymptomatic lady was diagnosed to have left main coronary artery atresia during the autopsy. We believe that patients with left main coronary artery atresia are the worst subset of AOCAPA and have a feeble LV function and severe MR at the time of diagnosis, with extensive collaterals. Surgical revascularisation using a bypass graft is reported for this subset of patients. However, in two of our patients, as the length of the bulbous left main coronary stump was adequate, with proximity to ascending aorta, we could transfer the left main coronary artery directly to the aorta. This technique is not reported in English medical literature to date (Fig. 7a–e).

For all our patients with ARCAPA, who were adolescent, asymptomatic and diagnosed elsewhere and referred to us, the dilemma existed regarding surgical intervention versus conservative management. Radke and colleagues [36], in a review, found that 40% of the patient were symptomatic and their study also confirmed the inability of the large dilated coronary arteries to appropriately fill the coronary sinus, owing to the steal phenomena with poor coronary flow reserve, until surgical correction. Mintz et al. [37] have also demonstrated the coronary steal phenomena in adult patients with ARCAPA. We believe that even if the patients are asymptomatic, they should be advised for surgical correction to establish two-coronary supply.

The literature search also revealed an extremely rare form of the anomalous origin of the single coronary artery from the PA, with successful surgical intervention in infancy [38]. We never came across such an anomaly in our series.

**Conclusion**

We believe and advocate that all cases of AOCAPA need surgical correction; otherwise, the patients will continue to have a progressively increasing left to right shunt and poor coronary flow, which will predispose to ischemia, arrhythmias and sudden death. Although echocardiography is diagnostic for most of these anomalies, there are other unusual presentations and variants, which may require definitive imaging using CT and magnetic resonance imaging. They are emerging as applicable diagnostic imaging modalities. One should always keep in mind the possibility of left main coronary atresia, ALCAPA and single coronary artery origin from the PA, as these are rare forms of AOCAPA. Nowadays, coronary reimplantation technique to re-establish the normal dual coronary supply has become the method of choice, with early mortality rate approaching to zero. LV dysfunction has been reported as the main risk factor for perioperative mortality. We routinely use prophylactic infusion of levosimendan in paediatric patients with impaired LV function. In patients operated in late infancy or childhood, LV function may not improve until the time of discharge, but gradual improvement in LV function and degree of MR is observed even after 7–8 years of surgery. We believe that the newborns and early infancy patients have better post-operative recovery and speedy recovery of their LV function. As the diastolic, longitudinal systolic function and LV torsion remain impaired, lifelong surveillance of ALCAPA patients is needed.

In our experience, in patients who had presented in late infancy and childhood with severe MR, the major cause of MR was ischemic lesions of the papillary muscles (organic MR). In this group, mitral annuloplasty, in association with coronary reimplantation, is indicated, due to the reduced likelihood of regression of MR after coronary revascularisation alone. The presence of left to right shunt provides accessory oxygenated blood supply to the LV and preserves the ventricular function. The presence of additional or accessory branch
from the RCA to the LV also helps in preserving the LV function. If there is sudden unexplained ventricular dysfunction in the postoperative patient with left to right shunt, anomalous origin of coronary artery from PA should be suspected. Early diagnosis and establishment of two coronary artery system can give the best outcome in this complex congenital condition.

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Compliance with ethical standards

Ethics approval and consent to participate Ethics approval was waived off being a retrospective study of a standard surgery. Informed consent was obtained from all patients.

Conflict of interest The author declares no competing interests.

Research involving human participants and/or animals This article does not contain any studies with animals performed by the author. Study was performed in accordance with the ethical standards as laid down in the 1964 Declaration of Helsinki and its later amendments or comparable ethical standards.

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