

Research progress in drug-coated balloon or/and directional atherectomy in treatment of complex femoropopliteal artery lesions

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[Abstract] Complex femoropopliteal artery lesions (CFPAL) is a clinical difficulty. Drug-coated balloon (DCB) can carry the drug directly to the lesion site for release, so that the drug can accumulate at the target site in a high concentration and inhibit intimal hyperplasia. Directional atherectomy (DA) can make the drug carried by DCB penetrate into the vascular wall better by removing the vascular plaque, and improve the clinical therapeutic effect. The application advances of DCB or/and DA in treatment of CFPAL were reviewed in this paper.

[Keywords] peripheral arterial disease; lower extremity; femoral artery; popliteal artery; arteriosclerosis obliterans; balloon; transluminal extraction-atherectomy therapy

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药物涂层球囊及定向斑块旋切术治疗复杂股腘动脉病变研究进展

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[摘要] 复杂股腘动脉病变(CFPAL)是临床治疗难点。药物涂层球囊(DCB)可携带药物直达病变部位释放,使药物在靶部位高浓度聚集,且可抑制内膜增生。定向斑块旋切术(DA)可通过切除血管斑块使DCB携带的药物更好地渗透入血管壁,提高临床治疗效果。本文对DCB、DA治疗CFPAL的应用进展进行综述。

[关键词] 外周血管疾病;下肢;股动脉;腘动脉;闭塞性动脉硬化;球囊;血管腔内斑块旋切术

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动脉硬化闭塞症(arteriosclerosis obliterans, ASO)是一种全身性疾病,主要累及大、中动脉,当病变累及腹主动脉下段及其远端主干动脉时,可引起下肢急性或慢性缺血。随着镍钛合金支架的出现^[1],血管腔内治疗逐渐成为下肢ASO临床首选血运重建方法;但术后血管或支架内再狭窄(in-stent restenosis, ISR)仍是困扰临床医师的主要问题,特别是复杂和特殊部位病变,如膝关节区,由于运动导致血管内产生持

续外压,可致ISR、支架断裂和闭塞发生率增加^[2]。药物涂层球囊(drug-coated balloon, DCB)可携带药物直达病变部位释放,同时抑制内膜增生。定向斑块旋切术(directional atherectomy, DA)可以通过切除血管斑块使DCB携带的药物更好地渗透入血管壁发挥临床治疗效果。DCB联合DA可显著改善血管腔内治疗ASO 1年后靶血管通畅率^[3-5]。本文对DCB和/或DA治疗复杂股腘动脉病变(complex

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femoropopliteal artery lesions, CFPAL)的研究进展进行综述。

1 CFPAL

2007年第2版泛大西洋国际共识(Transatlantic Inter-Society Consensus, TASC II)根据血管病变复杂程度将股腘动脉病变分为A~D型;其中C型和D型具有病变血管长、直径细、弥漫性钙化和多发血管性疾病等特点,加之部分患者伴有基础疾病(如糖尿病),发生二次狭窄、再闭塞和急性血栓形成等风险高^[6~7],并将此类下肢血管病变定义为CFPAL。有效治疗CFPAL是临床难点^[5]。我国2016版《下肢动脉硬化闭塞症诊治指南》推荐对TASC II A~C型股腘动脉病变首选腔内治疗,TASC II D型合并严重内科疾病或存在其他手术禁忌时也可选择腔内治疗^[8~9]。既往研究^[10~11]结果显示,单纯经皮腔内血管成形术(percutaneous transluminal angioplasty, PTA)或支架治疗CFPAL成功率较高,但远期通畅率不满意。近年来,DCB扩张术已成为治疗股腘动脉疾病的重要手段,相比单纯PTA治疗CFPAL更具优势。既往研究^[12~14]表明,DCB联合DA可有效降低长段靶病变血管重建率(target lesion revascularization, TLR)。

2 DA与DCB

目前国内多使用SilverHawk及TurboHawk斑块旋切系统,均为美国ev3公司产品,后者属于更新型,切割效率更高,适用于钙化较为严重的病变血管斑块。旋切系统主要由切割导管部分和驱动部分组成;为避免发生远端栓塞,可使用保护伞装置,其锥形头端可辅助扩张远端管腔并压迫斑块。该装置可有针对性地去除动脉粥样硬化斑块,并尽可能保留正常血管结构,缺点是需多次重复才能消除一定体积的斑块。Pantheris动脉斑块切除装置(美国Avenger公司)是最近获得美国食品药品监督管理局批准的新型DA装置。Pantheris导管配备光学相干断层成像(optical coherence tomography, OCT)装置,可在无电离辐射的情况下直接使动脉腔可视化,确保导管更安全、快速地到达病变血管部位,更有针对性地去除偏心斑块,增加旋切术的疗效和安全性,同时最大限度地降低非病变血管壁损伤风险,适用于治疗直径为3~7 mm的血管。此外,由OCT获取的数据可进行血管3D图像重建,使术者更好地观察病变部位,减少手术人员及患者X线暴露时间^[15~17]。

在以DCB行血管成形术之前先行DA治疗可使药物(如紫杉醇)更好地渗透入动脉壁,促进药物摄取

和利用,最大限度地避免切除顽固斑块所致局部炎症,降低新生内膜过度增生风险^[18~19]。目前在外周血管床上使用DCB存在两个主要问题:一是血流限制性夹层风险高,增加以球囊扩张支架进行补救治疗的概率;二是DCB治疗严重钙化血管病变预后较差^[13]。研究^[20~21]报道,单纯PTA治疗支架使用率为12.6%。另一项亚组分析^[22]结果显示,对于DCB血管成形术后的夹层,即使不植入支架,也不会导致不良结果。TEPE等^[13]发现,对于钙化病变,在抗增殖治疗前使用减容装置进行血管准备,可能对DCB治疗更有益。

3 DCB、DA治疗CFPAL

目前DCB主要用于治疗简单血管病变和TASC II A、B型病变。MICARI等^[23]认为DCB对于长段股浅动脉(superficial femoral artery, SFA)病变治疗效果较好,该研究采用DCB治疗105例存在SFA病变的CFPAL患者,病变血管平均长度为(251±71)mm,其中钙化病变占63.4%,完全闭塞占49.5%,病变的复杂性导致植入支架率相对较低(10.9%),随访12个月时血管通畅率为83.2%、TLR率仅4%,提示该法具有持续显著的临床效益,且踝肱指数(ankle-brachial index, ABI)、卢瑟福分级和生活质量评价结果均良好;但该研究属于非随机无对照临床试验,缺乏可比较性,需进一步完善。SHAMMAS等^[24]开展另一项单中心临床研究,对符合要求的75例新发或复发性股腘动脉病变患者进行回顾性分析,其中50例接受DA+普通PTA治疗、25例接受DA+DCB治疗,术后12个月2组免干预TLR率分别为68.0%、94.7%(P=0.002),16个月分别为54.0%、94.4%(P=0.002),表明DA+DCB的临床效果明显优于DA+普通PTA。CIOPPA等^[25]对30例有限制性跛行(n=18)及严重肢体缺血(n=12)患者(卢瑟福分级4.2±1.2)先给予DA治疗,随后进行DCB扩张,根据术中情况决定是否植入支架,手术成功率为100%,仅2例行补救性支架植入;术后随访1年,保肢率为100%,3例合并1型糖尿病患者出现再狭窄,需再次手术干预,术后一期通畅率>90%。

SCHMIDT等^[26]观察260例CFPAL患者共288处肢体病变,病变血管平均长度为(24.0±10.2)mm,平均ABI为0.56±0.22,严重肢体缺血发生率达26.4%,经DCB治疗后1年病变血管通畅率为79.2%,明显高于既往研究^[27]报道的22%~34%;但DCB治疗后随访2年,血管通畅率下降至53.7%,可能与手术操作的复杂性和患者本身合并多种基础疾病

有关。值得一提的是,该组对于 288 处病变中的 83 处在 DCB 血管成形术前进行血管腔内减容治疗(包括 DA、机械切除、激光消融),接受减容治疗的病变血管术后 1 年及 2 年平均通畅率分别为(85.3±4.3)% 和(64.0±6.1)%,相较于未接受减容治疗者[(76.8±3.2)% 和(49.6±4.0)%]有明显优势。

4 DCB、DA 治疗 ISR

传统 ISR 血管成形术后 6 个月血管开放率仅为 27%,收益期过短,效果欠佳^[28]。STABILE 等^[29]采用 DCB 治疗 39 处 SFA 病变后 ISR,术后随访 12 个月,一期通畅率为 92%。目前对于 DCB 治疗股腘动脉 ISR 的有效性和安全性仍存在争议。有研究^[30]对合并糖尿病 ISR 患者(多数患者出现危重肢体缺血,超过 1/2 患者存在 Tosaka III 级 ISR)分别给予紫杉醇 DCB 或单纯 PTA 治疗,术后 12 个月 DCB 组与单纯 PTA 组一期通畅率分别为 80.5% 及 28.2%,但第 3 年 2 组 TLR 发生率分别为 40% 和 43%,提示 DCB 治疗 ISR 并无远期优势。现有研究^[31]表明,单纯采用包括 DA 在内的血管腔内减容治疗 ISR 并不能获得较好的通畅率。对于 DCB 单独或联合 DA 治疗外周动脉 ISR 疗效尚待观察。

综上所述,DCB 单独或联合 DA 治疗 CFPAL 的安全性及有效性均较好,但其对于更复杂的腔内病变(如 ISR)的可行性及疗效仍需进一步探索。

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消息

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