Patellar groove replacement in patellar luxation with severe femoro-patellar osteoarthritis

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Summary
Objective: To report a novel method of treating femoro-patellar instability in association with severe femoro-patellar osteoarthritis, by substituting the femoral trochlear with a patellar groove replacement prosthesis.

Study design: Retrospective case series.

Methods: Preoperative lameness was scored from 0–4, and radiographic studies including standard positions for patellar luxation were obtained for evidence of malalignment and femoro-patellar osteoarthritis. Cases with or without previous surgeries were included. The size of trochlear implant was determined by transparent templates and confirmed intra-operatively with trials. Radiographic images, together with clinical examinations, were reviewed immediately and at three months postoperatively and at longer term when available.

Results: Thirty-five cases of patellar luxation ranging from grades II to IV were included. Eleven of these cases had prior surgical interventions which failed to stabilize the patella. Fourteen dogs required additional surgical procedures in conjunction with patellar groove replacement. Complications occurred in six patients, of which three required revision. Complete resolution of subjectively-assessed lameness was evident in 24/35 cases by the third month and in another seven of 35 patients on the longer term re-evaluations.

Clinical significance: Use of a patellar groove replacement prosthesis has the potential to decrease the lameness associated with severe femoro-patellar arthritis, to improve patellar stability, and to correct the alignment of the extensor mechanism.

Materials and methods

Medical records from two different institutions (Chirurgische Überweisungspraxis, Austria; Clinica Veterinaria Vezzoni srl, Italy) were reviewed to identify dogs affected by persistent patellar luxation and treated by patellar groove replacement. The inclusion criteria of the study were as follows: dogs affected with chronic medial or lateral patellar luxation, with or without cranial cruciate ligament injury, with intra-operative assessment of severe femoro-patellar osteoarthritis which would impede efficient trochleoplasty, and a minimum clinical and radiographic follow-up time period of three months. Exclusion criteria included severe femoro-tibial osteoarthritis requiring total knee replacement prosthesis.

Preoperative clinical assessment

Information obtained from the medical records included: date of presentation, date of surgery, signalment, clinical history, direction of patellar luxation (medial or lateral), grade of patellar luxation according to the Putnam scale of grade I to IV, previous surgical procedures, visual gait analysis by the surgeons (DL, AV) for the lameness associated with femoro-patellar osteoarthritis.
Implant design
The patellar groove replacement prosthesis was a two-component implant comprising of a base plate and a trochlear prosthesis (Figure 2). The base plate was perforated grade 4 titanium. To promote osseous integration it was coated with a glow discharge anodisation with incorporation of calcium phosphate. The trochlear prosthesis was made of grade 5 titanium (Ti6Al4V). It was anatomically shaped on the upper face, highly polished, and treated with amorphous diamond-like carbon coating to provide a very low coefficient of friction, scratch resistant surface. The base plate was provided with three conical holes matched to the conical feet of the trochlear prosthesis and was secured to the bone by either two bone screws (1.5 mm, titanium alloy) for the smaller sizes of the prosthesis, or by four bone screws (2.4 mm, titanium alloy) for the larger sizes. The feet of the trochlear prosthesis were aligned with the receiving holes of the base plate and were firmly seated by light tapping by hammer to achieve a press-fit locking. A gap of about 1 mm remained between the upper component and the base plate to ensure the conically fitted feet fully engaged the base plate.

Radiographic study and planning
Preoperative radiographs were obtained to document the patellar instability, patellar location, and to assess extensor alignment, and trochlear depth. The skyline view was used to document trochlear shape and depth. The cranio-caudal view and axial view of the femur plus the caudo-cranial view of the tibia, instead of the standard medio-lateral view of both femur and tibia, were used to document skeletal alignment.

From the medio-lateral radiographs, the size of the trochlear implant was determined using a transparent template over the radiograph (Figure 3), such that the base of the patellar groove replacement implant was aligned to a line from the point of the origin of the long digital extensor to the most proximal edge of the trochlea.

Surgical technique
All surgical procedures were performed by two board certified surgeons (AV and DL). Preoperatively, each dog was evaluated for any haematological or serum biochemical alterations. Animals were pre-medicated with morphine hydrochloride (0.2 mg/kg) and medetomidine hydrochloride (2 mcg/kg) and anaesthesia was induced with propofol (1-6 mg/kg) and maintained with isoflurane in oxygen. Cefazolin natricum (22 mg/kg IV) was administered at the time of induction and repeated after 90 minutes. Dogs were positioned in dorsal recumbency and aseptically prepared for surgery. Either a medial parapatellar approach for medial patellar luxation or a lateral parapatellar approach for lateral patellar luxation was performed (8). After reflection of the patella, an osteotomy of the trochlea, parallel to the frontal plane was performed with an oscillating saw starting distally at the level of the origin of the tendon of the long digital extensor muscle, ending proximally at the proximal end of the trochlea. In the case of femoral torsion, the osteotomy was slightly tilted medial or lateral to correct for torsion, keeping 25° to 35° as a normal reference for femoral neck anteversion angle (9). In medial patellar lu-
and stability of the reduction was assessed with flexion and extension of the stifle. The necessary adjustments made included insertion of a larger or smaller trial prosthesis or removal of an additional slice of bone. In addition, a proper fit of the patella inside the prosthetic trial trochlea was verified. Adjustments were made by finding the best medio-lateral and proximo-distal position of the trial prosthesis to stabilize the patella. Tibial tuberosity transposition was not generally required, since the patella could be realigned by simply moving the troclear prosthesis medially or laterally in the frontal plane. Moreover, in cases of medial patellar luxation, the prosthetic groove could be fixed more proximally to counteract the tendency of medial luxation occurring with extension of the stifle, while in cases of lateral patellar luxation it could be fixed more distally to counteract the tendency of lateral luxation happening during stifle flexion (11). Femoral varus or valgus was partially compensated by rotating the prosthesis in the frontal plane. Once the preferred position of the trial prosthesis was found, the bone was gently marked with an osteotome so that the base plate of the final patellar groove replacement prosthesis could be fixed in the same position (Figure 5, Figure 6). The base plate was stabilized to the bone with two to four titanium cortical screws, depending on the size of the implant. The final prosthesis was press-fitted to the base plate by aligning the three conical pegs of the prosthetic to the corresponding holes in the base plate and gently tapping the prosthetic with a Teflon hammer. The patella was then reduced and tested again in flexion and extension. Some patelloplasty was performed if required by trimming the lateral and medial borders of the patella if it was too wide or flat to seat properly inside the trochlea. Capsulorraphy with imbrication of the opposite side to the luxation was performed, and leaving the capsule open on the same side of the luxation in the grade IV cases. Subcutis and skin were routinely sutured completing the surgical procedure.

Additional surgical procedures

When the underlying limb deformities were too severe and it was not possible to
realign the quadriceps mechanism with patellar groove replacement alone, or when cruciate ligament failure was concomitant, additional surgical procedures were performed in combination with patellar groove replacement including tibial tuberosity transposition, distal femoral osteotomy, proximal tibial osteotomy, tibial plateau levelling osteotomy or tibial tuberosity advancement (Appendix Table 1: Available online at www.vcot-online.com).

**Postoperative patient evaluation and management**

Postoperative cranio-caudal and mediolateral radiographs of the full femur with the stifle positioned in flexion and extension were obtained to evaluate the osteotomy site, and to confirm the patellar reduction and prosthesis position. When femoral torsion was compensated with a tilted osteotomy, the axial radiographic view was also taken. A modified Robert Jones bandage was applied for the first 24 – 48 hours postoperatively. Pain was controlled during the first two days with administration of morphine hydrochloride (0.2 mg/kg SC q 6 hr) and meloxicam (0.1 mg/kg SC q 24 hr) for five days. Cephadroxil (20 mg/kg PO q 12 hr) was administered for five days. Postoperative exercise restriction to leash walking was enforced for two months, which was followed by a gradual increase in the following month up to normal levels of activity. When indicated in chronic conditions with reduced range-of-motion of the affected stifle, physiotherapy was recommended.

**Follow-up**

During the follow-up period, outcome assessment was focused on clinical and radiographic examination at three months after surgery, and more frequently and at longer term when indicated or possible. Radiographic assessments included stability and position of the patellar groove replacement implant, patellar reduction, any alteration of bone interface with the prosthesis, and progression of osteoarthritis. Clinical evaluation assessed the status of extensor realignment, passive range-of-motion with or without signs of pain, muscle tone, and strength. Gait evaluation assessed the absence or the degree of lameness. This analysis was divided into a short-term follow-up (3 months) and into a long-term follow-up (6 months and over) when available (Figure 7).

**Statistical analysis**

A paired t-test or non-parametric signed rank test was used to assess the difference in lameness scores. The test statistic of Shapiro-Wilk was used to compare continuous measurements to a normal distribution. A p-value of <0.05 was used to determine statistical significance. Commercially available software was used for the statistical analysis.

**Results**

**Preoperative clinical findings and surgical details**

Findings are summarized in Appendix Table 1 (Available online at www.vcot-online.com). Of the 35 cases that met the inclusion criteria, 16 were males, one was a neutered male, 12 were females, and six were female-spayed. Of the 35 cases, medial patellar luxation was present in 29, of which 22 were grade III luxation and six

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h Metacam: Boehringer Ingelheim Vetmedica, Bracknell, UK
i Cefa Cure: Intervet Int, Boxmeer, The Netherlands
j SAS version 9.4: SAS Institute, Cary, NC, USA

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Figure 5 Fixation of the base plate and insertion of the trochlæar prosthesis in a plastic bone model. Image courtesy of KYON AG (PGR Brochure).

Figure 6 Case 9: Osteotomy of the degenerated trochlea, fixation of the base plate and insertion of the trochlæar prosthesis.
were a grade IV. Lateral patellar luxation was present in six cases, of which one was a grade II luxation and five were a grade IV. The age ranged from eight to 125 months with a mean of 54.5 months. Body weight ranged from 1.5 – 97.5 kg with a mean of 21.9 kg. All 35 cases had short-term (3 months) follow-up, whereas 33/35 had a long-term follow-up (6 to 56 months). The median time for long-term follow-up was 12 months and the mean 19 months.

The sizes of prosthesis used in these cases were as follows: 1 (n = 3), 2 (n = 3), 3 (n = 4), 4 (n = 5), 5 (n = 1), 6 (n = 2), 7 (n = 8), 8 (n = 2), 9 (n = 5), 10 (n = 2).

Outcome

Findings are summarized in Appendix Table 1 (Available online at www.vcot-online.com). All 35 cases were lame prior to surgery with lameness scores ranging from 2 – 4 (median score of 3). Three months following surgery, all cases improved with lameness scores ranging from 0 to 1 (mean score of 0.3). Specifically 24/35 cases were not lame at three months while 11/35 cases were still lame but with a lesser score compared to the preoperative lameness. The long-term evaluation determined that 23 of the 33 with a lameness score of 0 at three months remained unchanged in longer term evaluation, seven improved from a lameness score of 1 to 0 and three remained with a score of 1. Differences in lameness scores between the short-term and long-term follow-up periods were also significant (p <0.0156) indicating a continued improvement beyond three months following patellar groove replacement. The null hypothesis was rejected with these results, confirming a positive clinical effect with the patellar groove replacement procedure.

Not all cases had patellar groove replacement surgery alone as 14 patients required additional surgical treatment including tibial tuberosity transposition (n = 5), tibial plateau levelling osteotomy (n = 4), tibial tuberosity advancement (n = 1), distal femoral osteotomy (n = 3), and proximal tibial osteotomy (n = 3), and a lateral fabellar-tibial suture in one case. However in 21 cases where the underlying limb deformities were less severe, patellar groove replacement was the only procedure done.

Complications

Complications were observed in six cases, of which three required surgical revision. In one unrevised case (case 2), an oversized implant resulted in capsule tension and thickening. Two others were a Chihuahua (case 4) and a Yorkshire Terrier (case 33) with medial tilting of the patella inside the patellar groove replacement prosthesis which nevertheless did not cause lameness or discomfort on stifle manipulation. Of the three cases requiring surgical revision, a mongrel (case 7) demonstrated painful impingement during range-of-motion in the third and fourth week after surgery, where the patella slipped over the distal end of the prosthesis creating a “clunk” sensation. Revision performed one month after surgery included excision of 3–4 mm of excess bone over the extensor fossa and of redundant fibrous tissue from the proximal border of the patella, which led to complete improvement at three months. A large breed dog (case 20) experienced sudden lameness four months after surgery due to dislodgement of the prosthesis. The case was successfully revised with re-fitting of a replacement prosthetic trochlea including the base which had been altered by friction with the loose implant. The dog had regained normal function by eight months following revision. A medium size dog (case 22) had a recurrence of medial patellar luxation one month after surgery requiring limb alignment because of excessive distal femoral varus (anatomical lateral distal femoral angle 107°) and external femoral torsion (femoral antversion angle 15°). Distal femoral osteotomy with a final anatomic lateral distal femoral angle of 95° and femoral anteverversion angle of 30° was performed. Patellar reduction was confirmed 11 months following the additional osteotomies. At three months following the patellar groove replacement procedure, three dogs (cases 3, 5, and 19) that had chronic patellar luxation and severe preoperative osteoarthritis had some progression of the arthritis evident on the follow-up radiographs. In the remaining 32 cases progression of osteoarthritis was not observed.

Discussion

The patellar groove replacement procedure has the potential to restore function to dogs with severe femoro-patellar arthritis. The patellar groove replacement also provides...
for improved patellar coverage and alignment of the extensor mechanism with or without tibial tuberosity transposition or other corrective osteotomy. In our opinion, balanced tracking of the patella is more likely to result with the patellar groove replacement procedure than with plain or resection trochleoplasty because the tall ridges and smooth contours of the prosthesis allow for more deep gliding of the patella. The smooth surface of the prosthesis also provides a low friction surface for smooth gliding of the patella, reducing tissue damage and inflammation. Furthermore, moderate varus-valgus and external-internal torsion of the femur can be compensated by setting the prosthesis in a calculated angle of inclination in the frontal plane.

Reduction of lameness assessed by clinical survey suggests a prompt and notable postoperative recovery. Lameness scores dropped to 0 during the first few weeks in 24 dogs, and in further seven cases in the longer term. The three dogs with persistent grade 1 lameness in the long-term also had concurrent cruciate ligament rupture with the chronic patellar luxation, which could explain that outcome. Multiple preoperative and adjunctive surgical procedures may increase the likelihood of residual morbidity. The mild progression of osteoarthritis observed in three cases with chronic luxation and severe preoperative osteoarthritis did not correlate with clinical performance.

Oversizing the prosthesis could lead to soft tissue impingement suggesting that proper planning and intra-operative testing with a trial prosthesis is necessary to find the right size for each patient. Two dogs (cases 4 and 33) had medial patellar tilt visible on the skyline radiographic views despite a more medial implant position, probably because of a proportionally wider patellar groove replacement prosthesis size to patellar size and a persistent malalignment of the extensor mechanism requiring tibial tuberosity transposition.

While the patellar groove replacement prosthesis has the potential to improve function in case of severe femoro-patellar arthritis, it cannot be expected to compensate alone for marked bone deformities occurring in cases of grade III and IV patellar luxation. Case 22 demonstrated the requirement for additional corrective femoral osteotomy for a successful restoration of function. In this case, a recurrence of patellar luxation occurred after patellar groove replacement because the femoral deformity and subsequent malalignment were overlooked. However, where the underlying limb deformities were less severe, fixation of the patellar groove replacement prosthesis in line with the quadriceps mechanism allowed proper realignment without further surgical procedures. Also, the capability of fixing the patellar groove replacement prosthesis slightly more proximally in the case of medial patellar luxation or more distally in the case of lateral patellar luxation than the original trochlea, allows treatment of patella alta or baja. In this current series of cases, patellar groove replacement was performed concurrently with other surgical procedures in 14 cases. In eight of these cases, the additional surgery was done to correct severe femoral or tibial deformities or both. In the other six cases, patellar groove replacement was performed with procedures designed to address cranial cruciate deficiency.

Bone ingrowth and absence of any adverse bone reaction are characteristics of medical grade titanium (12, 13). The patellar groove replacement prosthesis is a titanium implant designed for bone ingrowth and minimal tissue reaction. Aseptic loosening was not recognized in this series of clinical cases, suggesting appropriate bone integration with the base plate. Actual ingrowth cannot be confirmed by radiographic imaging and clinical performance alone, but considering the long-term stability and function of the implant evaluated in the 21 cases with a minimum of one year follow-up indicates successful bone tissue ingrowth and absence of inflammatory tissue reaction.

Case 20 had a dislocation of the trochlear prosthesis which may have been due to an incomplete press fit at the time of implantation. Loosening of the base plate from the underlying bone was not evident in this case; nevertheless dislodgment of the prosthesis constitutes an incidence of implant loosening. Dislodgment of the trochlear prosthesis could potentially also occur because of patellar mal-tracking when patellar alignment was not addressed with specific osteotomies if required.

The incidence of complications decreased with an increasing number of cases, suggesting that surgeon experience played a role in the complication rate. Experience improved the selection of the appropriate size of implant preventing oversizing. Proper positioning of the implant also improved with the experience of the surgeon.

A primary limitation of this study was the survey basis as the principal assessment of outcome. Surveys of the surgeons were done with a base standardized system to improve reliability. However, the authors recognize the inherent bias that exists in such data and a repeated study with improved design including a control group and an evaluation by a clinician who is blinded to the operative procedure would be indicated.

Nevertheless, this study has shown that the patellar groove replacement implant is well tolerated without adverse reactions. The patellar groove replacement procedure has the potential to improve function in cases of severe femoro-patellar osteoarthritides due to chronic patellar luxation, and it has the potential to improve the alignment of the extensor mechanism in the case of moderate malalignment and keeping the patella permanently reduced. More extensive clinical studies with confirmed statistical outcomes will be essential in establishing the role of the patellar groove replacement implant in the treatment of femoro-patellar abnormalities in the dog. Moreover, computed tomographic evaluation of the skeletal abnormalities causing patellar luxation could provide more precise measurements for surgical planning compared to plain radiographs.

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Conflict of interest

A. Vezzoni teaches the PGR technique at KYON courses; no payments however are received from KYON for using this technique in clinical cases. No other conflicts of interest to declare.

References