Liver transplant remains the definitive treatment for end-stage liver disease. In 2017, 182 Canadians either died while on the wait list or became too sick to undergo the operation.¹ A CMAJ article by Shemie highlighted the current trend toward increasing use of donation after circulatory death (DCD) liver grafts. However, these marginal grafts are prone to higher complication rates, particularly biliary complications. In addition, many procured DCD livers are then deemed unsuitable for transplant. Despite these limitations, DCD grafts represent an important resource to address the current organ shortage, and as such there are research efforts directed toward improving the use of and outcomes for transplantation of these grafts. We review the current progress in DCD liver transplantation.

Reducing wait list mortality among patients awaiting liver transplantation remains a substantial challenge because of organ shortage. In efforts to expand the donor pool there has been a trend toward increased use of donation after circulatory death (DCD) liver grafts. However, these marginal grafts are prone to higher complication rates, particularly biliary complications. In addition, many procured DCD livers are then deemed unsuitable for transplant. Despite these limitations, DCD grafts represent an important resource to address the current organ shortage, and as such there are research efforts directed toward improving the use of and outcomes for transplantation of these grafts. We review the current progress in DCD liver transplantation.

La réduction du nombre de personnes en attente d’une greffe de foie qui décèdent avant la transplantation demeure un défi important en raison de la pénurie d’organes. On remarque actuellement une tendance à la hausse dans l’utilisation de greffons de foie provenant de don après décès circulatoire (DDC) dans le but d’élargir le bassin de donneurs. Ces greffons marginaux sont toutefois associés à des taux de complications plus élevés, particulièrement pour ce qui est des complications biliaires. De plus, de nombreux foies obtenus à la suite d’un DDC sont jugés inadmissibles à la greffe. Malgré ces restrictions, les greffons provenant de DDC représentent une importante ressource pour atténuer la pénurie d’organes. Des initiatives de recherche sont donc actuellement en cours dans le but d’améliorer leur taux d’utilisation et les issues des transplantations. Nous analyisons ici l’état actuel des progrès pour les transplantations de foie provenant de DDC.

Donation after circulatory death differs from the more common donation after neurological death (DND). Before procurement, DCD livers are exposed to a period of warm ischemia time (WIT) after cardiopulmonary arrest in which the organ is no longer perfused; in DND livers, perfusion persists right up until the moment of procurement in the operating room. As a result, higher complication rates persist.³ Although DCD liver transplants have increased in recent years, in 2017 15% of the DCD liver grafts procured in Canada were subsequently discarded before transplantation.³ The high discard rate of the grafts and higher complication rates induced by WIT suggest that the current preservation strategy of static cold storage (SCS) is not well tolerated by these injured grafts. We present a brief overview of DCD liver transplant outcomes and review the progress in improving DCD outcomes (e.g., donor selection, use of thrombolytic medications, machine perfusion preservation strategies).
Literature search

We searched PubMed using the Medical Subject Headings “donation after circulatory death” and “liver transplant.” This search yielded 211 articles. Of these, we reviewed those that matched best. Relevant articles from the references listed in select articles were also reviewed. When possible, we discussed the highest level of evidence in randomized controlled trials; however, this level of evidence was limited and, therefore, observational data were also reviewed.

Current outcomes in DCD liver transplantation

A 2011 meta-analysis that included 489 DCD and 4455 DND recipients revealed a 1.6-fold increase in 1-year mortality and a 2.1-fold increased risk of 1-year graft failure for DCD grafts compared with DND.4 More recently, however, a meta-analysis including more than 12 000 patients found no difference in patient or graft survival when comparing DCD and DND liver transplants.5

The same improvement has not been observed regarding ischemic cholangiopathy (IC), defined as the occurrence of multiple intrahepatic strictures in the absence of hepatic artery thrombosis or stenosis.6 Ischemic cholangiopathy is 2.5 times more likely in DCD liver grafts than DND.5 While hepatocytes receive dual blood supply from both the portal vein and hepatic artery, biliary cells receive only arterial blood.7 Both DND and DCD livers demonstrate loss of intraluminal biliary epithelium as a result of procurement and preservation; however, only those with significant damage to the deep peribiliary plexus, the site of progenitor cells for regeneration of the biliary ducts, show an increased risk of IC.8 Thus it has been hypothesized that IC develops from an impaired ability to regenerate biliary cells after revascularization secondary to peribiliary vascular plexus injury in DCD grafts.9

Ischemic cholangiopathy leads to higher rates of graft failure, longer hospital admissions, increased biliary procedures, retransplantation and overall increased health care costs.8 Improving IC rates in DCD transplants remains a substantial challenge for ongoing research efforts.

Donor factors affecting DCD liver transplant outcomes

Donor age

Older donor age has been identified consistently as a risk factor for complications in DCD liver transplantation.10 Donor age older than 50 years has been shown to predict risk for IC11 and graft failure.10 However, when other risk factors were minimized, Schlegel and colleagues found no difference in biliary or overall complications in DCD liver grafts between cohorts older or younger than 60 years.12 They concluded that older DCD donors may provide viable grafts when other risk factors are mitigated.

Donor obesity

Rising obesity rates have resulted in increasing numbers of donors with steatosis.13 In DCD liver grafts from obese donors, there has been poor tolerance of SCS evidenced by worse transplant outcomes.13 Elevated body mass index (BMI) has been associated with increased IC as well as worse graft and patient survival.10–12,14 Liver steatosis leads to lower adenosine triphosphate (ATP) levels, microcirculatory dysfunction, increased inflammation and reactive oxygen species production with ultimately more severe ischemia reperfusion injury.13 Improving outcomes of this expanding subset of donor livers has become an area of keen research interest for machine perfusion.

Warm ischemia time

For DCD procurement, the WIT cut-off in many centres is 30 minutes. Both graft failure and IC have been linked to prolonged WIT.10,14,15 Tun-Abraham and colleagues16 evaluated their single-centre series, dividing their cohort into early (July 2006 to June 2011) and late (July 2011 to July 2016) periods. The late group showed reduced times from incision to arterial cannulation and organ flush.16 This was accompanied by a statistically significant reduction in IC for the late group.16 These improvements were attributed to a learning curve, suggesting procurements done by staff experienced in DCD could improve outcomes.16 In addition a meta-analysis looking at DCD liver transplant outcomes between those who had life support withdrawn in the intensive care unit compared with the operating room found that both graft survival and rate of IC were improved by withdrawal of life support in the operating room.17 This varies by institution; however, a shift in policies to allow withdrawal of life support in the operating room may contribute to improved outcomes.

Cold ischemia time

Cold ischemia time (CIT) has been identified as a significant risk factor for poor outcomes in DCD liver transplantation.18,19 In order to reduce the risk of IC and graft failure, it has been suggested that CIT be limited to less than 10 hours.18 Others suggest even more strict time limits of less than 9 hours of total ischemia time.11

Thrombolytic medications during DCD transplantation

The addition of thrombolytic medications has received recent research interest (Table 1). Tissue plasminogen activator (tPA) was injected via the hepatic artery on the back
table based on the hypothesis that microthrombi from stasis during WIT led to damaged peribiliary vasculature and ultimately contributed to IC.\textsuperscript{20} In this initial study, 22 DCD transplants were carried out with the tPA protocol; 2 (9\%) recipients developed IC, which is lower than previously reported; however, excessive bleeding was encountered in 14 (64\%) of the 22 recipients.\textsuperscript{20} Subsequent studies modified the tPA protocol and injected the drug via the hepatic artery after venous reperfusion in the recipient.\textsuperscript{21,24} A description of the approaches for tPA administration in DCD liver transplantation has been described in detail elsewhere.\textsuperscript{25} The more recent studies using tPA showed improved 1- and 3-year patient and graft survival and a significant reduction in IC relative to controls.\textsuperscript{21,24} These studies did not show the same increased transfusion requirement previously described by Hashimoto and colleagues.\textsuperscript{20} A detailed meta-analysis that included 249 patients in the tPA group and 178 patients not receiving tPA showed a significant reduction in IC and retransplant rates for the tPA group without increased transfusion requirements, leading the authors to conclude that tPA provides an advantage for preventing IC.\textsuperscript{21} Further supporting the use of tPA protocols for DCD liver grafts is the recent demonstration of cost savings with this intervention.\textsuperscript{26}

**PROGRESS IN EX-SITU MACHINE PERFUSION**

The higher incidence of biliary complications and graft failure with DCD grafts has been attributed to the damage incurred during WIT and subsequent SCS. Increased research interest in machine perfusion has led to the clinical use of this modality as an alternative preservation strategy (Table 2). Machine perfusion is classified broadly by the temperature of the perfusate: hypothermic machine perfusion (HMP), subnormothermic machine perfusion (SNMP), and normothermic machine perfusion (NMP).

**Hypothermic machine perfusion**

Providing oxygen in the perfusate at subphysiologic flow rates and temperatures allows for improved mitochondrial and endothelial protection.\textsuperscript{16} In a large animal model, DCD livers undergoing HMP showed lower alanine aminotransferase (ALT) and aspartate aminotransferase (AST) levels,\textsuperscript{17} and improved preservation of the biliary microvasculature relative to SCS.\textsuperscript{38} There were also significant improvements in ATP levels, bile production, bile composition and lactate levels relative to those transplanted after SCS.\textsuperscript{28,39} Several clinical studies using HMP have now been completed (Table 2), the largest of which compared 50 DCD livers treated with HMP to 50 DCD SCS livers and 50 DND controls. There was a significantly lower rate of IC relative to the DCD SCS group with increased survival and reduced retransplant rate at 1 year.\textsuperscript{27} At 5-year follow-up, graft survival was 94\% compared with 78\% in DCD grafts without HMP.\textsuperscript{20}

A criticism of this modality is that it remains difficult to assess liver function during perfusion, as metabolic

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### Table 1. Thrombolytic medications in DCD liver transplantation

| Study            | No. of DCD patients | Thrombolytic protocol | Thrombolytic dose | Summary of key outcomes                                |
|------------------|---------------------|-----------------------|-------------------|---------------------------------------------------------|
| Hashimoto et al.\textsuperscript{20} | n = 22              | tPA injected in the hepatic artery on the back table | tPA protocol dose (n = 12): 0.6 mg/100 g graft weight | Graft survival at 1 and 3 years was 81\%. Ischemic cholangiopathy developed in 9\% while 64\% had excessive bleeding. |
|                  |                     |                       | tPA reduced dose for high risk of bleeding (n = 10): 0.2–0.4 mg/100 g graft weight |                                      |
| Seal et al.\textsuperscript{21} | tPA n = 85, no tPA n = 33 | Centre specific: tPA injected into the hepatic artery at the time of portal vein anastomosis or at the time of hepatic artery anastomosis | Centre specific: tPA 2 mg or 0.5 mg/kg donor weight | tPA use improved 1- and 3-year patient and graft survival, reduced ischemic cholangiopathy and did not increase the risk of bleeding. |
| Pietersen et al.\textsuperscript{22} | Urokinase n = 63 (17 DCD), No urokinase n = 122 (28 DCD) | Urokinase injected into hepatic artery on back table immediately before implantation | Urokinase 250 000 IU | Urokinase use did not lower ischemic cholangiopathy rates. There was no difference in graft survival and no increase in blood transfusion requirements. |
| Kubal et al.\textsuperscript{23} | tPA n = 30, no tPA n = 61 (historic controls) | tPA injected into hepatic artery: tPA 100 mg mixed with 1 L normal saline initial flush (900 mL), then additional 100 mL of tPA solution following organ flush with HTK | tPA total dose 100 mg | No difference in overall survival. tPA flush resulted in a significant reduction in ischemic cholangiopathy without increased bleeding risk. |
| Bohorquez et al.\textsuperscript{24} | tPA n = 100, no tPA n = 38 (historic controls) | tPA injected into hepatic artery immediately after portal vein reperfusion | tPA 2 mg | tPA use improved 1- and 3-year graft survival. There was no statistically significant difference in ischemic cholangiopathy and no increased bleeding risk. |

DCD = donation after circulatory death; HTK = histidine–tryptophan–ketoglutarate; tPA = tissue plasminogen activator.
| Study | No. of patients | Perfusion temperature | Median WIT, min | Median CIT, hr | Perfusion duration, hr | Outcome measures | Key findings |
|-------|----------------|----------------------|----------------|---------------|----------------------|-----------------|--------------|
| Dutkowski et al.\textsuperscript{27} | HMP: 25 DCD SCS: 50 DCD historic controls | HMP | 36 | 6.6 | 2 | AST, ALT, bilirubin, INR, early allograft dysfunction, ischemic cholangiopathy, hospital/ICU stay, graft loss, 1-year graft survival | HMP-treated livers had a significant decrease in peak ALT, ischemic cholangiopathy and improved 1-year graft survival. |
| van Rijn et al.\textsuperscript{28} | HMP: 10 DCD SCS: 20 DCD historic controls | HMP | 26 | 6 | 2.1 | 6-month graft survival, 1-year graft and patient survival, technical safety of perfusion, microbiological testing, postoperative complications, lactate, ALT, ALP, GGT, prothrombin time, bilirubin, ICU/hospital length of stay, ischemic cholangiopathy, ATP | Significant reduction in postoperative ALT, ALP, GGT, bilirubin in HMP group. No significant difference in hospital/ICU length of stay. Nonsignificant reduction in ischemic cholangiopathy in HMP group. |
| van Rijn et al.\textsuperscript{29} (same patient population as above study) | HMP: 10 DCD SCS: 20 DCD historic controls | HMP | 26 | 6 | 2.1 | Biliary histologic injury, nonanastomotic biliary strictures | HMP resulted in decreased ischemia reperfusion injury of the bile ducts with improved preservation of the peribiliary glands. |
| Schlegel et al.\textsuperscript{30} | HMP: 50 DCD SCS: 50 DCD historic controls SCS: 50 DND historic controls | HMP | 31 | — | — | INR, ALT, hospital/ICU length of stay, postoperative complications, ischemic cholangiopathy | Significantly improved 5-year graft survival in HMP group. HMP group also had less ischemic cholangiopathy but with longer hospital stays. |
| Ravikumar et al.\textsuperscript{31} | NMP: 20 (4 DCD) SCS: 40 (8 DCD) historic controls | NMP | 21 | 8.9 | 9.3 | 30-day graft survival, AST, ALP, bilirubin, INR, patient and graft survival, 6-month graft function, length of stay | Lower peak AST in the first 7 postoperative days in the NMP group. No difference in graft or patient survival. No difference in length of ICU or hospital stay. |
| Selzner et al.\textsuperscript{32} | NMP: 10 (2 DCD) SCS: 30 (6 DCD) historic controls | NMP | 49 | 9.7 | 8 | 90-day patient and graft survival, AST, ALT, INR, bilirubin, creatinine, need for dialysis, length of stay, complications | No difference in transaminase levels, patient or graft survival, or length of hospital/ICU stay. |
| Bral et al.\textsuperscript{33} | NMP: 9 (4 DCD) SCS: 30 (8 DCD) historic controls | NMP | 21.5 | 3.8 | 11.5 | 30-day graft and patient survival, AST, bilirubin, INR, 6-month graft and patient survival, 6-month biliary complications | No difference in 30-day or 6-month graft survival. No ischemic cholangiopathy in NMP group. NMP group had significantly longer ICU and hospital length of stay |
| Nasralla et al.\textsuperscript{34} | NMP 170 (63 DCD) SCS 164 (60 DCD) RCT | NMP | 21 | 7.8 | 9.1 | Peak AST, organ discard rate, post-reperfusion syndrome, primary nonfunction, early allograft dysfunction, length of hospital/ICU stay, need for renal replacement therapy, 6-month cholangiopathy, 1-year patient and graft survival | NMP group showed lower peak AST, less early allograft dysfunction and lower discard rates relative to SCS. These findings were more pronounced in the DCD subgroup. There was no statistically significant difference in biliary outcomes, hospital/ICU stay or patient and graft survival. |
| Bral et al.\textsuperscript{35} | NMP: 17 (4 DCD) NMP following SCS: 26 (6 DCD) | NMP | 20 SCS + NMP: 21 | NMP: 3.2 SCS + NMP: 6 | 8.1 | 90-day and 6-month patient and graft survival, AST, ALT, ALP, bilirubin, INR, early allograft dysfunction, 6-month biliary and vascular complications | No difference in graft or patient survival. Biliary complications between groups were equivalent. |

ALP = alkaline phosphatase; ALT = alanine aminotransferase; AST = aspartate aminotransferase; CIT = cold ischemia time; DCD = donation after circulatory death; DND = donation after neurological death; GGT = γ-glutamyl transferase; HMP = hypothermic machine perfusion; ICU = intensive care unit; INR = international normalized ratio; NMP = normothermic machine perfusion; RCT = randomized controlled trial; SCS = static cold storage; WIT = warm ischemia time.
activity is minimized under hypothermic conditions. Thus, defining criteria of organs suitable for transplantation will be an important goal if this modality is to be used more widely in clinical practice. Hypothermic machine perfusion has shown benefit following periods of SCS, which in the Canadian population, where organs are being transported long distances, would allow for implementation without substantial alteration of current procurement strategies. An ongoing randomized trial (NCT02584283) will shed further light on the potential of this technology.

**Subnormothermic machine perfusion and controlled oxygenated rewarming**

In animal studies, controlled oxygenated rewarming (COR) with gradual rewarming followed by perfusing at subnormothermic temperatures (20–21°C) was superior to HMP and SCS, with improved serum enzyme levels, portal vein resistance, bile production and histological injury scores. It also demonstrated improved ATP recovery relative to NMP or SCS. The first clinical series involved 6 patients who underwent COR for 90 minutes following SCS and were compared with a historic cohort. Controlled oxygenated rewarming had a statistically significant reduction in peak AST, and at 6-month follow-up all patients had normal liver function tests. However this series did not include any DCD grafts. There are limited data regarding SNMP or COR for DCD liver grafts. In a series of perfusions consisting of 5 discarded DCD human livers, SNMP demonstrated increased ATP levels and clearance of lactate during 3 hours of SNMP. More evidence is required in order to make any conclusions regarding the use of SNMP/COR for DCD liver grafts.

**Normothermic machine perfusion**

Normothermic machine perfusion aims to recreate physiologic conditions to recover and assess liver grafts before transplantation. Large animal models have demonstrated lower transaminase levels, improved lactate clearance and better histologic preservation of hepatocytes relative to SCS. In addition, NMP has shown improved preservation of biliary endothelial regeneration capacity, which is hypothesized to prevent the development of IC. It is also being investigated as a modality to “de-fat” livers from obese donors. However, consensus on the optimal NMP perfusate composition, circuit set-up, and viability criteria have not yet been reached. Nonetheless, following the positive results of animal studies, multiple clinical trials have now been completed (Table 2). In the first phase-1 clinical trial, 20 livers (4 DCD) were preserved using NMP and compared with 40 (8 DCD) controls preserved with SCS. The NMP group demonstrated significantly lower median peak AST levels in the first 7 days post-transplant (417 IU [interquartile range (IQR) 84–4681]) relative to the SCS controls (4902 [IQR 218–8786], \( p = 0.03 \)), with 100% 30-day graft survival in the NMP group. These results led the authors to conclude that this technology was safe and had the potential to increase the number of quality transplantable livers. This study was followed up by the publication of 2 North American phase-1 trials similar in design to that of Ravikumar and colleagues. Selzner and colleagues compared 10 livers (2 DCD) preserved with NMP to 30 (6 DCD) liver grafts preserved with SCS, and Bral and colleagues compared 9 grafts (4 DCD) preserved with NMP to 30 (8 DCD) grafts preserved with SCS. Though neither of these 2 studies was able to show the same statistically significant reduction in transaminase levels, there was no difference in graft survival or ischemic cholangiopathy. The study by Bral and colleagues demonstrated higher overall transaminase levels than the other 2 phase-1 clinical trials, which was attributed to there being a greater proportion of DCD grafts in their study. There was also a statistically significant increase in the median length of stay in the intensive care unit (ICU) for the NMP group relative to SCS controls (16 d \( v. \) 4 d, \( p = 0.004 \)). Taken together, these phase-1 trials established the safety and feasibility of NMP leading to commencement of the first randomized control trial. Given the low numbers of DCD grafts in these initial trials and the results of Bral and colleagues, the capacity to improve the quality of DCD grafts using NMP remains to be determined.

In the first randomized control trial, Nasralla and colleagues compared 10 livers (2 DCD) preserved with 164 livers (64 DCD) that were randomized to NMP and SCS, respectively. There was no significant difference in 1-year graft or patient survival; however, there was a statistically significant reduction in peak AST in the NMP group compared with the SCS group (4902 [IQR 218–8786], \( p = 0.03 \)). Taken together, these phase-1 trials established the safety and feasibility of NMP leading to commencement of the first randomized control trial. Given the low numbers of DCD grafts in these initial trials and the results of Bral and colleagues, the capacity to improve the quality of DCD grafts using NMP remains to be determined.

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unclear whether any interventional strategies will ultimately mitigate the risk of IC.

These initial published studies using NMP have perfusion initiated at the time of procurement. This requires the donor procurement either to take place in the transplant centre with the perfusion device, or that the perfusion device be portable. Bral and colleagues have recently completed the first clinical study investigating the use of NMP following a period of SCS. Normothermic machine perfusion was initiated after SCS in 26 livers (6 DCD) and compared with 17 grafts (4 DCD) that had NMP initiated at the time of procurement. There was no significant difference in graft or patient survival or transaminase levels between groups. Biliary complications were also similar between groups.

While this study included only a small number of patients, particularly for DCF grafts, it suggests that NMP can be used safely after a period of SCS without compromising outcomes. This will require further study with larger numbers of patients before being implemented more widely in clinical practice; however, as the authors point out, in Canada many procurements require air travel and therefore being able to maintain current procurement practices and then use NMP after arrival back to the transplant centre would greatly simplify the logistics of implementing this technology.

**CONCLUSION**

Donation after circulatory death liver transplantation has resulted in an increase in transplants being performed in Canada; however, discard rates are substantial, and higher rates of IC persist. Donor selection remains critical, while thrombolytic protocols have shown early benefits for graft survival and IC. Machine perfusion shows promise in both increasing use and improving outcomes; however, clinical data are still emerging, particularly for DCD grafts. Ongoing research aims to optimize machine perfusion protocols, establish reliable viability criteria and demonstrate consistent long-term outcomes before widespread clinical implementation.

**Affiliations:** From the Department of Surgery, Division of General Surgery, University of Alberta Hospital, Edmonton, Alta. (Nostedt, Shapiro, Bigam); the Department of Physiology, University of Alberta, Edmonton, Alta. (Freed); and the Department of Surgery, Division of Cardiac Surgery, University of Alberta, Alberta Heart Institute, Edmonton, Alta. (Freed).

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