Use of minimal invasive extracorporeal circulation in cardiac surgery: principles, definitions and potential benefits. A position paper from the Minimal invasive Extra-Corporeal Technologies international Society (MiECTiS)


a Cardiothoracic Department, AHEPA University Hospital, Thessaloniki, Greece
b Department of Anesthesiology and Perioperative Medicine, University of Western Ontario, London, Canada
c Department of Cardiothoracic Surgery, MediClin Heart Centre Coswig, Coswig, Germany
d Department of Anaesthesia and Intensive Care, Policlinico S. Donato, Milan, Italy
e Department of Cardiovascular Surgery, University of Bern, Bern, Switzerland
f University Pierre and Marie Curie (Paris 06), Paris, France
g Department of Cardiac Surgery, Regensburg, Germany
h Department of Anesthesiology and Pain Therapy, University of Bern, Bern, Switzerland
i Department of Cardiothoracic Surgery, Hammersmith Hospital, London, UK
j Department of Cardiothoracic and Vascular Surgery, Ulm University, Ulm, Germany
k Department of Cardiothoracic Surgery, Amsterdam Medical Center, Amsterdam, Netherlands
l Heart Centre, University Hospital Ghent, Gent, Belgium
m Department of Thoracic and Cardiovascular Surgery, Braunschweig, Germany
n Department of Adult Cardiac Surgery, Mater Dei Hospital, Bari, Italy
o Department of Cardiac Surgery, Oulu University Hospital, Oulu, Finland
p Department of Surgery, Sidra Medical & Research Centre, Doha, Qatar
q Department of Cardiothoracic Surgery, Maastricht University Medical Centre, Maastricht, Netherlands
r Department of Thoracic and Cardiovascular Surgery, University Hospital of the Rhine University Bochum, Bad Oeynhausen, Germany
s University Health Network, Toronto, Canada
t St Antonius Hospital, Nieuwegein, Netherlands
u Department of Cardiovascular Surgery, Medline Hospitals, Adana, Turkey
v Department of Cardiothoracic Surgery, Wessex Cardiac Centre, University Hospital Southampton, Hampshire, UK
w Department of Cardiac Surgery, Bristol Heart Institute, Bristol, UK
x Department of Cardiothoracic Surgery, German Heart Centre, Berlin, Germany

* Corresponding author. Clinic for Cardiovascular Surgery, University Hospital Bern and University of Bern 3010 Bern, Switzerland. Tel: +41-31-6322375; fax: +41-31-6324443; e-mail: thierry.carrel@insel.ch (T. Carrel).

Received 11 September 2015; received in revised form 17 November 2015; accepted 25 November 2015

Summary

Minimal invasive extracorporeal circulation (MiECC) systems have initiated important efforts within science and technology to further improve the biocompatibility of cardiopulmonary bypass components to minimize the adverse effects and improve end-organ protection. The Minimal invasive Extra-Corporeal Technologies international Society was founded to create an international forum for the exchange of ideas on clinical application and research of minimal invasive extracorporeal circulation technology. The present work is a consensus document developed to standardize the terminology and the definition of minimal invasive extracorporeal circulation technology as well as to provide recommendations for the clinical practice. The goal of this manuscript is to promote the use of MiECC systems into clinical practice as a multidisciplinary strategy involving cardiac surgeons, anaesthesiologists and perfusionists.

Keywords: Extracorporeal circulation • Minimally invasive extracorporeal circulation • Cardiopulmonary bypass • Modular systems • Systemic inflammation reaction syndrome • Complications
INTRODUCTION

Substantial experience has been accumulated with cardiac procedures performed using extracorporeal circulation (ECC) over the last decades. Several technological improvements have been realized, thus making cardiopulmonary bypass (CPB) the gold standard equipment for the majority of cardiac surgical procedures. This has contributed to improved perioperative and long-term results, despite an increasing prevalence of elderly and high-risk patients [1]. For the most frequent procedure, coronary artery bypass grafting (CABG), CPB provides optimal conditions (bloodless field and arrested heart) to allow the most complete myocardial revascularization and additionally offers the possibility to perform other procedures such as valve repair or replacement and aortic surgery [2].

Major drawbacks of CPB are the adverse systemic effects triggered by a systemic inflammatory response syndrome (SIRS), which is mainly caused by the contact of blood with air and foreign surfaces [3,4]. Trials have shown that the inflammatory response to CPB adversely influences clinical outcome [5,6] although CPB cannot be considered as the main cause of postoperative morbidity.

Since the beginning of extracorporeal perfusion, the main inputs have been focused on one objective—to reduce the adverse effects of CPB. Perfusionists and bioengineers have developed optimized ‘CPB systems’ that combined the best features derived from perfusion science. The idea was to create a system that integrates all modifications into one combined set-up, known as the minimally invasive extracorporeal circulation (MiECC) system [7]. This concept has further initiated important new efforts to improve the biocompatibility of CPB components and minimize the side-effects.

Despite clinical advantages that have been reported in several papers [8], the penetration of MiECC technology into clinical practice remains extremely low. There is also significant heterogeneity between the various systems. Low implementation of MiECC may be due to the inability to predict which aspects of MiECC are beneficial, because several elements may act both interactively and/or independently, e.g. coated surfaces, closed systems, anticoagulation strategies, shed blood separation and reduced priming volumes.

The Minimal invasive Extra-Corporeal Technologies international Society (MIECTiS) was founded to create an international forum to exchange ideas on clinical practice and research in the field of minimally invasive extracorporeal circulation technology (www.miectis.org). The Society brings together, under a scientific interdisciplinary association, cardiac surgeons, anaesthesiologists, perfusionists and basic researchers.

The present work is a consensus document developed to standardize the terminology around MiECC technology and to provide recommendations for clinical practice. The authors have graded the levels of evidence and classified the findings listed below using the criteria recommended by the American Heart Association and the American College of Cardiology Task Force on Practice Guidelines (Table 1). The authors represent a multidisciplinary group to promote evidence-based perfusion practice to improve clinical outcomes.

METHODS

The initiative to analyse the current practice was based on a questionnaire that was written by the Steering Committee of MIECTiS (Kyriakos Anastasiadis, Adrian Bauer, Thierry Carrel, Erich Gygax, John Murkin, Marco Ranucci, Jan Schaarschmidt). During an Expert Consensus Meeting, the statements were discussed and, subsequently, this consensus paper was developed. For each statement, the best available published evidence derived from meta-analyses of peer-reviewed literature, randomized controlled trials (RCTs) and data coming from large cohort studies were considered. Relevant studies were searched in PubMed (1975–present), Embase (January 1980–present) and Cochrane review of aggregate data for reports written in any language. The full PubMed search strategy is available in Supplementary Table 1.

Moreover, hand or computerized search involving the recent (1999–2014) conference proceedings from the Society of Thoracic Surgeons, European Association for Cardiothoracic Surgery and European Society for Cardiovascular Surgery and the American Association for Thoracic Surgery Annual Meetings was performed; ClinicalTrials.gov was explored in order to identify any ongoing or unpublished trials (Table 2).

Recommendations and evidence-based practice guidelines

Expert Committee statements are presented in Table 3. Evidence-based clinical practice guidelines are presented in Table 4.

Terminology

MiECC refers to a combined strategy of surgical approach, anaesthesiological and perfusion management and is not limited to the CPB circuit alone.

Several terms have been used to describe a MiECC circuit: miniaturized extracorporeal circulation (MECC), mini-extracorporeal...
<table>
<thead>
<tr>
<th>Author, journal, date [Ref.]</th>
<th>Study type</th>
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<th>Key results</th>
<th>Comments</th>
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<tbody>
<tr>
<td>Wiesenack et al., Artif Organs, 2004 [10]</td>
<td>Retrospective analysis</td>
<td>CABG</td>
<td>485 MiECC/485 CCPB</td>
<td>Type I</td>
<td>Higher MAP and mean pump flow rate during in MiECC. Reduced frequency of vasoactive drug administration in MiECC patients ($P &lt; 0.05$). Maximum values of lactate concentration during bypass were significantly higher in CCPB. Minimum values of haemoglobin as an indicator of haemodilution were higher in MiECC patients ($P &lt; 0.05$). Transfusion of packed red blood cells during surgery and during the complete perioperative course was significantly larger in CCPB ($P &lt; 0.05$). 30-day mortality was similar between groups. Incidence of postoperative complications was significantly higher in CCPB ($P &lt; 0.05$).</td>
<td>First reported large series showing improved perfusion characteristics and clinical results</td>
</tr>
<tr>
<td>Yilmaz et al., Interact CardioVasc Thorac Surg, 2010 [11]</td>
<td>Prospective cohort study</td>
<td>CABG + AVR</td>
<td>65 MiECC/135 CCPB</td>
<td>Type III</td>
<td>Reduced preoperative haemoglobin drop and higher haemoglobin at discharge in MiECC ($P = 0.03$). Reduced blood products requirements in MiECC ($P = 0.004$). No differences were noted in pulmonary complications, neurological events or mortality.</td>
<td>Feasibility study</td>
</tr>
<tr>
<td>Anastasiadis et al., Perfusion, 2015 [12]</td>
<td>Prospective cohort study</td>
<td>Various cardiac case-mix</td>
<td>50 consecutive patients</td>
<td>Type IV</td>
<td>Technical success 100%. 4% conversion rate from Type III to Type IV (modular MiECC). Clinical study on modular type IV MiECC in all types of cardiac surgery (feasibility and safety study).</td>
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<tr>
<td>El-Essawi et al., Perfusion, 2011 [13]</td>
<td>Multicentre RCT (six centres)</td>
<td>CABG and/or AVR</td>
<td>252 MiECC/248 CCPB</td>
<td>Type IV</td>
<td>No operative mortality or device-related complications. Cardiomyotmy suction was necessitated by major bleeding in 10 patients. Integration of a hard-shell reservoir was deemed necessary for air handling in 1 patient. Transfusion requirement ($P = 0.001$), incidence of atrial fibrillation ($P = 0.03$) and the incidence of major adverse events ($P = 0.02$) were all in favour of the MiECC group. No thromboembolic events in either group. Low-dose group had lower 24-h mean postoperative blood loss ($P = 0.001$) and reduced rate of transfusion of allogeneic blood ($P = 0.01$). Four patients in the control group received a total of 10 units of packed red blood cells and, in the low-dose group, no transfusions were given ($P = 0.046$). No patient was reoperated because of bleeding. ICU stay was significantly shorter in the low-dose group ($P = 0.020$). Patients in low-dose group were less dependent on oxygen on the first postoperative day ($P = 0.034$), better mobilized ($P = 0.006$) and had less pain ($P = 0.019$).</td>
<td>Focus on modular type IV MiECC in CABG and/or AVR</td>
</tr>
<tr>
<td>Fromes et al., Anaesthesia, 2011 [14]</td>
<td>Retrospective analysis</td>
<td>CABG</td>
<td>100 patients 300 IU/kg heparin/68 patients 145 IU/kg heparin</td>
<td>Type II</td>
<td>No thromboembolic events in either group. Low-dose group had lower 24-h mean postoperative blood loss ($P = 0.001$) and reduced rate of transfusion of allogeneic blood ($P = 0.01$). Implementation of low-dose heparin protocol</td>
<td></td>
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<tr>
<td>Nilsson et al., Interact CardioVasc Thorac Surg, 2012 [15]</td>
<td>RCT</td>
<td>CABG</td>
<td>27 low-dose heparin/29 regular dose</td>
<td>Type II</td>
<td>Four patients in the control group received a total of 10 units of packed red blood cells and, in the low-dose group, no transfusions were given ($P = 0.046$). No patient was reoperated because of bleeding. ICU stay was significantly shorter in the low-dose group ($P = 0.020$). Patients in low-dose group were less dependent on oxygen on the first postoperative day ($P = 0.034$), better mobilized ($P = 0.006$) and had less pain ($P = 0.019$).</td>
<td>Feasibility of low-dose heparin</td>
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### Table 2: (Continued)

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<thead>
<tr>
<th>Author, journal, date</th>
<th>Study type</th>
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<th>Type of MiECC circuit</th>
<th>Key results</th>
<th>Comments</th>
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<tbody>
<tr>
<td>Anastasiadis et al., J Cardiothorac Vasc Anesth, 2013 [16]</td>
<td>RCT</td>
<td>CABG</td>
<td>60 MiECC/60 CCPB</td>
<td>Type II</td>
<td>Incidence of fast-track recovery was significantly higher in patients undergoing MiECC ($P = 0.006$). MiECC was recognized as a strong independent predictor of early recovery ($P = 0.011$). Duration of mechanical ventilation and cardiac recovery unit stay were significantly lower in patients undergoing MiECC. Need for blood transfusion, duration of inotropic support, need for intra-aortic balloon pump, development of postoperative atrial fibrillation and renal failure were significantly lower in patients undergoing MiECC.</td>
<td>Focus on fast-track protocols</td>
</tr>
<tr>
<td>Anastasiadis et al., Perfusion, 2010 [17]</td>
<td>RCT</td>
<td>CABG</td>
<td>50 MiECC/49 CCPB</td>
<td>Type I</td>
<td>Less haemodilution ($P = 0.001$), markedly less haemolysis ($P &lt; 0.001$) and better preservation of the coagulation system integrity ($P = 0.01$) favouring MiECC group. Less bank blood requirements were noted and a quicker recovery, as far as mechanical ventilation support and ICU stay are concerned, in MiECC group.</td>
<td>Focus on haematological effects</td>
</tr>
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<td>Haneya et al., ASAIO J, 2013 [18]</td>
<td>Retrospective cohort analysis</td>
<td>CABG</td>
<td>1073 MiECC/872 CCPB</td>
<td>Type I</td>
<td>Postoperative creatine kinase and lactate levels were significantly lower in the MiECC group ($P &lt; 0.001$). No difference in postoperative blood loss between the groups. Intraoperative and postoperative transfusion requirements were significantly lower in the MiECC group ($P &lt; 0.05$). MiECC patients had lower incidences of postoperative acute renal failure, low cardiac output syndrome, shorter intensive care unit lengths of stay and reduced 30-day mortality ($P &lt; 0.05$).</td>
<td>Focus on patients with preoperative anaemia</td>
</tr>
<tr>
<td>Zangrillo et al., J Thorac Cardiovasc Surg, 2010 [19]</td>
<td>Meta-analysis (16 RCTs)</td>
<td>CABG or AVR</td>
<td>803 MiECC/816 CCPB</td>
<td>MiECC was associated with significant reductions of neurological damage ($P = 0.008$), reduction in peak cardiac troponin ($P &lt; 0.001$), and in the number of transfused patients ($P &lt; 0.001$).</td>
<td>No difference in mortality was noted.</td>
<td>Meta-analysis</td>
</tr>
<tr>
<td>Anastasiadis et al., Int J Cardiol, 2013 [20]</td>
<td>Meta-analysis (24 RCTs)</td>
<td>CABG or AVR</td>
<td>1387 MiECC/1383 CCPB</td>
<td>MiECC was associated with a significant decrease in mortality ($P = 0.02$), in the risk of postoperative myocardial infarction ($P = 0.03$) and reduced rate of neurological events ($P = 0.08$).</td>
<td>MiECC was associated with significantly reduced systemic inflammatory response, haemodilution, need for red blood cell transfusion, reduced levels of peak troponin release, incidence of low cardiac output syndrome, need for inotropic support, peak creatinine level, occurrence of postoperative atrial fibrillation, duration of mechanical ventilation and ICU stay.</td>
<td>The largest meta-analysis</td>
</tr>
<tr>
<td>Rahe-Meyer et al., Artif Organs, 2010 [21]</td>
<td>Prospective cohort study</td>
<td>CABG</td>
<td>44 MiECC/44 CCPB</td>
<td>Type I</td>
<td>Aggregation decreased significantly in both groups as early as 30 min after the institution of CPB ($P &lt; 0.05$) and recovered within the first 24 h postoperatively, without reaching the preoperative level. Intraoperative aggregometry values reflected a significantly more severe reduction of platelet function in CCPB group ($P &lt; 0.01$).</td>
<td>Focus on coagulation</td>
</tr>
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<td>Study</td>
<td>Design</td>
<td>Patients</td>
<td>Procedures</td>
<td>Cases</td>
<td>Type</td>
<td>Key Findings</td>
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<tr>
<td>El-Essawi et al., Perfusion, 2013 [22]</td>
<td>Cohort</td>
<td>Various</td>
<td>29 patients</td>
<td>22 CABG ± AVR</td>
<td>Type IV</td>
<td>Mean decrease in haemoglobin was 2.1 ± 1.3 g/dl during cardiopulmonary bypass and 3.4 ± 1.4 g/dl at discharge. Lowest postoperative haemoglobin level was 9.3 ± 1.8 g/dl. MiECC system allowed a reduced haemodilution (P &lt; 0.05). Mononuclear phagocytes dropped in a more important manner in CCPB group (P = 0.002). No significant release of IL-1b was observed in either group. By the end of CPB, IL-6 levels were significantly lower in MiECC group (P = 0.04), despite a higher monocyte count. Plasma levels of TNF-α increased significantly in CCPB group (P = 0.002). Neutrophil elastase release was significantly reduced in MiECC group (P = 0.001). Platelet count remained at higher values with MiECC β-Thromboglobulin levels showed slightly lower platelet activation in the MiECC group (P = 0.10). Focus on SIRS</td>
</tr>
<tr>
<td>Fromes et al., Eur J Cardiothorac Surg, 2002 [23]</td>
<td>RCT</td>
<td>CABG</td>
<td>22 CABG ± AVR</td>
<td>7 various case-mix</td>
<td>Type I</td>
<td>Mean decrease in haemoglobin was 2.1 ± 1.3 g/dl during cardiopulmonary bypass and 3.4 ± 1.4 g/dl at discharge. Lowest postoperative haemoglobin level was 9.3 ± 1.8 g/dl. MiECC system allowed a reduced haemodilution (P &lt; 0.05). Mononuclear phagocytes dropped in a more important manner in CCPB group (P = 0.002). No significant release of IL-1b was observed in either group. By the end of CPB, IL-6 levels were significantly lower in MiECC group (P = 0.04), despite a higher monocyte count. Plasma levels of TNF-α increased significantly in CCPB group (P = 0.002). Neutrophil elastase release was significantly reduced in MiECC group (P = 0.001). Platelet count remained at higher values with MiECC β-Thromboglobulin levels showed slightly lower platelet activation in the MiECC group (P = 0.10). Focus on SIRS</td>
</tr>
<tr>
<td>Immer et al., Ann Thorac Surg, 2007 [24]</td>
<td>Cohort</td>
<td>CABG</td>
<td>1053 MiECC/353 CCPB</td>
<td>Type I + smart suction device</td>
<td>TnI was significantly lower in the MiECC group (P &lt; 0.05). Incidence of AF was significantly reduced in MiECC (P &lt; 0.05). Inflammatory markers (IL-6, SC5b-9) were lower in MiECC patients (P = 0.05). Propensity score analysis confirmed faster recovery in MiECC patients and lower incidence of AF.</td>
<td>Feasibility/safety study</td>
</tr>
<tr>
<td>Abdel-Rahman et al., Ann Thorac Surg, 2005 [25]</td>
<td>RCT</td>
<td>CABG</td>
<td>101 MiECC/103 CCPB</td>
<td>Type II</td>
<td>Intraoperative blood loss was significantly higher in CCPB group (P &lt; 0.0001) as well as the need of fresh frozen plasma. Postoperative chest drainage did not differ significantly between groups. One hour after CPB, PMNE as well as TCC were significantly lower in MiECC group (P &lt; 0.001). Platelet count remained at higher values with MiECC β-Thromboglobulin levels showed slightly lower platelet activation in the MiECC group (P = 0.10). Focus on SIRS</td>
<td>Feasibility/safety study</td>
</tr>
<tr>
<td>Ohata et al., J Artif Organs, 2007 [26]</td>
<td>RCT</td>
<td>CABG</td>
<td>15 MiECC/15 CCPB</td>
<td>Type I</td>
<td>Neutrophil elastase levels were lower in MiECC group at PODs 1 and 2 (P = 0.013). IL-8 levels were reduced in MiECC patients on POD 1 (P = 0.016). Intraoperative blood loss and transfusion volumes were significantly lower in MiECC group (P = 0.012). Number of distal anastomoses was lowest in the OPCABG group, but comparable for MiECC and CCPB patients. Postoperative ventilation time, release of creatinine kinase, catecholamine therapy, drainage loss and transfusion requirements were lower in the MiECC and OPCABG groups, whereas stay in the ICU was shorter only in the latter (P &lt; 0.05). Focus on SIRS</td>
<td>Feasibility/safety study</td>
</tr>
<tr>
<td>Puehler et al., Ann Thorac Surg, 2009 [27]</td>
<td>Cohort</td>
<td>CABG</td>
<td>558 MiECC/558 CCPB/558 OPCAB</td>
<td>Type I</td>
<td>In-hospital mortality for elective and urgent/emergent patients was lower in the MiECC and OPCAB groups (P &lt; 0.05). Number of distal anastomoses was lowest in the OPCABG group, but comparable for MiECC and CCPB patients. Postoperative ventilation time, release of creatinine kinase, catecholamine therapy, drainage loss and transfusion requirements were lower in the MiECC and OPCABG groups, whereas stay in the ICU was shorter only in the latter (P &lt; 0.05). Focus on SIRS</td>
<td>Feasibility/safety study</td>
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<tr>
<td>Biancani, Heart, 2009 [28]</td>
<td>Meta-analysis (13 RCTs)</td>
<td>CABG or AVR</td>
<td>562 MiECC/599 CCPB</td>
<td>MiECC was associated with reduced mortality during the immediate postoperative period, not reaching statistical significance (P = 0.25). Postoperative stroke rate was significantly lower in MiECC group (P = 0.05). Length of ICU stay was similar in both groups (P = 0.87). MiECC was associated with a significantly lower amount of postoperative blood loss (P = 0.0002) along with a higher platelet count 6 h after surgery (P = 0.03).</td>
<td>Meta-analysis</td>
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<tr>
<td>Author, journal, date [Ref.]</td>
<td>Study type</td>
<td>Type of procedure</td>
<td>Patient groups</td>
<td>Type of MiECC circuit</td>
<td>Key results</td>
<td>Comments</td>
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<td>Liebold, J Thorac Cardiovasc Surg, 2006 [29]</td>
<td>RCT</td>
<td>CABG</td>
<td>20 MiECC/20 CCPB</td>
<td>Type I</td>
<td>CCPB group showed a highly significant reduction in both cerebral oxygenated haemoglobin and tissue oxygenation index from the start to the end of cardiopulmonary bypass ($P &lt; 0.01$). The rate of decrease in cerebral oxygenated haemoglobin after aortic cannulation was faster in the CCPB group ($P &lt; 0.001$). No significant changes with respect to cerebral oxygenated haemoglobin or tissue oxygenation index occurred MiECC group, except at the beginning of rewarming ($P &lt; 0.01$). Total embolic count, as well as gaseous embolic count, in the left and right median cerebral arteries was significantly lower in MiECC group (all $P &lt; 0.05$). Postoperative bleeding was greater ($P &lt; 0.05$) and the transfusion rate was higher ($P &lt; 0.05$) in CCPB group.</td>
<td>Focus on cerebral protection</td>
</tr>
<tr>
<td>Zanatta, J Cardiothorac Vasc Anesth, 2013 [30]</td>
<td>Retrospective cohort</td>
<td>CABG</td>
<td>19 MiECC (CABG)/18 CCPB (AVR or MVR)/18 port-access MVR</td>
<td>Type I</td>
<td>The number of solid microemboli and gaseous microemboli was significantly reduced in MiECC group ($P &lt; 0.001$).</td>
<td>Focus on cerebral protection</td>
</tr>
<tr>
<td>Camboni, ASAIO J, 2009 [31]</td>
<td>RCT</td>
<td>CABG</td>
<td>42 MiECC type I/10 MiECC type II/41 CCPB</td>
<td>Type I and II</td>
<td>MiECC resulted in reduced microbubble activity compared with CCPB ($P = 0.02$). Postoperative neuropsychological dysfunction ($P = 0.45$), renal dysfunction ($P = 0.67$), days of hospitalization ($P = 0.27$) and 30-day mortality ($P = 0.30$) did not differ between groups.</td>
<td>Focus on cerebral protection</td>
</tr>
<tr>
<td>Anastasiadis et al., Heart, 2011 [32]</td>
<td>RCT</td>
<td>CABG</td>
<td>29 MiECC/31 CCPB</td>
<td>Type I</td>
<td>MiECC was associated with improved cerebral perfusion during CPB. Less patients operated on with MiECC experienced at least one episode of cerebral desaturation ($P = 0.04$) with similar duration. At discharge, patients operated on with MiECC showed a significantly improved performance on complex scanning, visual tracking, focused attention and long-term memory. At 3 months, significantly improved performance was also evident on visuospatial perception, executive function, verbal working memory and short-term memory. Patients operated on with MiECC experienced a significantly lower risk of early cognitive decline both at discharge ($P = 0.03$) and at 3-month evaluation ($P &lt; 0.01$).</td>
<td>Focus on neurocognitive outcome</td>
</tr>
<tr>
<td>Reineke et al., Interact CardioVasc Thorac Surg, 2014 [33]</td>
<td>Cohort study</td>
<td>CABG</td>
<td>31 MiECC</td>
<td>Type I + smart suction device</td>
<td>MiECC does not adversely affect cognitive brain function after CABG.</td>
<td>Focus on neurocognitive outcome</td>
</tr>
<tr>
<td>Study</td>
<td>Design</td>
<td>Procedure</td>
<td>MI ECC</td>
<td>CCPB</td>
<td>Type</td>
<td>Summary</td>
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<tr>
<td>Gynaydin, Perfusion, 2009 [34]</td>
<td>RCT</td>
<td>CABG</td>
<td>20 MiECC/20 CCPB</td>
<td>Type IV</td>
<td>Serum IL-6 levels were significantly lower in the MiECC group ($P &lt; 0.05$). C3a levels were significantly less in the MiECC group ($P &lt; 0.01$). CK-MB levels in coronary sinus blood demonstrated well-preserved myocardium in the MiECC group. Percentage expression of neutrophil CD11b/CD18 levels was significantly lower in the MiECC group ($P &lt; 0.05$). No significant differences in air handling characteristics or free plasma haemoglobin levels in either circuit. rSO2 measurements were significantly better in the MiECC group ($P &lt; 0.05$). Blood protein adsorption analysis of oxygenator membranes demonstrated a significantly increased amount of microalbumin on CCPB fibres ($P &lt; 0.05$).</td>
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<tr>
<td>Bennett et al., Perfusion, 2014 [35]</td>
<td>Cohort study</td>
<td>CABG and/or AVR</td>
<td>39 MiECC 41 CCPB</td>
<td>Type II</td>
<td>The average indexed bypass pump flow was significantly lower with MiECC with same average oxygen delivery. Patients in the CCPB group had a greater duration and severity of cerebral desaturation, which was significantly associated with low flows during CPB, whereas desaturation with MiECC was associated with low perioperative haemoglobin concentration.</td>
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<tr>
<td>Panday et al., Interact Cardiovasc Thorac Surg, 2009 [36]</td>
<td>Prospective cohort study</td>
<td>CABG</td>
<td>220 MiECC 1143 CCPB 109 OPCAB</td>
<td>Type II</td>
<td>Operative mortality rates were comparable in all three groups. The mean number of distal anastomoses was higher in MiECC and CCPB groups than OPCAB group ($P = 0.01$). Arrhythmia occurred in 25% of the MiECC group, in 35.6% of the CCPB group ($P = 0.05$) and in 21.7% of the OPCAB group. 3% of the MiECC group suffered neurocognitive disorders perioperatively compared with 7% of the CCPB group ($P = 0.05$) and 3% of the OPCAB group. The median number of blood transfusions per patient was lower in MiECC and OPCAB groups ($P &lt; 0.0001$).</td>
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<tr>
<td>Remadi, Am Heart J, 2006 [37]</td>
<td>RCT</td>
<td>CABG</td>
<td>200 MiECC/200 CCPB</td>
<td>Type I + suction device</td>
<td>Operative mortality rate similar between groups. Low cardiac output syndrome was reduced in MiECC group ($P &lt; 0.001$). Inflammatory response was significantly reduced in MiECC. C-reactive protein release postoperatively was significantly higher in CCPB group. Significantly higher decrease of haematocrit and haemoglobin rate in CCPB group. Intraoperative transfusion rate was reduced in MiECC group ($P &lt; 0.0001$). Patients in the CCPB group had significantly higher levels of postoperative blood creatinine and urea. MiECC exerts beneficial haemodynamic effects but does not prevent AKI. Fewer patients developed a decline in eGFR &lt; 60 mL/min/1.73 m² in MiECC ($P &lt; 0.001$). The incidence of eGFR decrease by &gt;50% did not differ ($P = 0.20$). Temporary dialysis was reduced in MiECC group ($P &lt; 0.001$). MiECC is renoprotective in the early postoperative period but cannot prevent AKI.</td>
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<tr>
<td>Diez et al., ASAIO J, 2009 [38]</td>
<td>Retrospective observational study</td>
<td>CABG</td>
<td>1685 MiECC /3046 CCPB</td>
<td>Type I</td>
<td>Focus on SIRS and haemodilution</td>
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<td>Focus on cerebral protection</td>
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<td>Focus on blood transfusion</td>
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<td>Feasibility/safety study</td>
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<td>Focus on renal function</td>
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<tr>
<td>Author et al., journal, date</td>
<td>Study type</td>
<td>Type of procedure</td>
<td>Patient groups</td>
<td>Type of MiECC circuit</td>
<td>Key results</td>
<td>Comments</td>
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<tr>
<td>Huybregts et al., Ann Thorac Surg, 2007</td>
<td>RCT</td>
<td>CABG</td>
<td>25 MiECC/24 CCPB</td>
<td>Type II</td>
<td>MiECC was associated with attenuation of on-pump haemodilution, improved haemostatic status with reduced platelet consumption and platelet activation, decreased postoperative bleeding and minimized transfusion requirements. MIoCC showed reduced leucocytosis and decreased urinary interleukin-6. Levels of urine NGAL were on average 3-fold lower and urinary intestinal fatty acid binding protein was 40% decreased in patients operated on MiECC.</td>
<td>Focus on renal and intestinal function</td>
</tr>
<tr>
<td>Capuano et al., Interact CardioVasc Thorac Surg, 2009</td>
<td>Prospective cohort study</td>
<td>CABG</td>
<td>30 MiECC/30 CCPB</td>
<td>Type II</td>
<td>CCPB group showed a significant NGAL concentration increase from preoperative during the first postoperative day (P &lt; 0.05). No patient in MiECC group developed AKI. Renal function is better protected during MiECC as demonstrated by NGAL levels.</td>
<td>Focus on renal injury</td>
</tr>
<tr>
<td>Benedetto et al., Ann Thorac Surg, 2009</td>
<td>Prospective cohort study</td>
<td>CABG</td>
<td>104 MiECC/601 CCPB</td>
<td>Type II</td>
<td>Overall incidence of AKI for patients undergoing MiECC was reduced (P = 0.03). MAP values were significantly higher in the MiECC group (P = 0.002). MiECC patients received significantly less norepinephrine (P = 0.045).</td>
<td>Focus on perfusion characteristics</td>
</tr>
<tr>
<td>Bauer et al., J Extra Corporeal Technol, 2010</td>
<td>RCT</td>
<td>CABG</td>
<td>18 MiECC/22 CCPB</td>
<td>Type II</td>
<td>MIoCC patients demonstrated significantly lower levels of TnT at 6, 12 and 24 h and CK-MB levels at 6 and 12 h. Markers of myocardial oxidative stress or activity were significantly lower in MiECC group compared with CCPB and OPCAB (P = 0.04 and 0.03, respectively).</td>
<td>Focus on myocardial protection</td>
</tr>
<tr>
<td>Skrabal, ASAIO J, 2007</td>
<td>RCT</td>
<td>CABG</td>
<td>30 MiECC/30 CCPB</td>
<td>Type I</td>
<td>MIoCC group showed significantly lower median TnT levels compared with CCPB and OPCAB (P &lt; 0.003). HFABP, IFABP and a-GST levels were significantly higher during CCPB compared with OPCAB and MiECC (P &lt; 0.009). There was a trend towards higher median CC16 levels in the CCPB group (P &lt; 0.07).</td>
<td>Focus on end-organ protection</td>
</tr>
<tr>
<td>Van Boven et al., Eur J Cardiothorac Surg, 2008</td>
<td>RCT</td>
<td>CABG</td>
<td>10 MiECC/10 CCP 10 OPCAB</td>
<td>Type I</td>
<td>Liver function as measured by disappearance rate of indocyanine green was markedly increased after cardiac surgery without significant differences between groups.</td>
<td>Focus on liver function</td>
</tr>
<tr>
<td>Van Boven et al., Eur J Anaesthesiol, 2013</td>
<td>RCT</td>
<td>CABG</td>
<td>20 MiECC/20 CCP 20 OPCAB</td>
<td>Type I</td>
<td>There is an impairment of microvascular perfusion during CCPB (P = 0.034). Changes in functional capillary density indicate a faster recovery of the microvascular perfusion in MiECC during the reperfusion period (P = 0.017).</td>
<td>Focus on microvascular perfusion</td>
</tr>
<tr>
<td>Prasser et al., Perfusion, 2007</td>
<td>RCT</td>
<td>CABG</td>
<td>10 MiECC/10 CCPB</td>
<td>Type I</td>
<td>The overall cardiac injury was significantly lower in the MiECC group as measured by TnT (P = 0.02).</td>
<td>Focus on myocardial protection</td>
</tr>
<tr>
<td>Donndorf et al., J Thorac Cardiovasc Surg, 2012</td>
<td>RCT</td>
<td>CABG</td>
<td>20 MiECC/20 CCPB</td>
<td>Type I</td>
<td>The overall cardiac injury was significantly lower in the MiECC group as measured by TnT (P = 0.02).</td>
<td>Focus on myocardial protection</td>
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<tr>
<td>Study</td>
<td>Design</td>
<td>Procedure</td>
<td>Patients</td>
<td>Outcome Measures</td>
<td>Results</td>
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<td>Haneya et al., Eur J Cardiothorac Surg, 2009 [49]</td>
<td>Retrospective cohort study</td>
<td>CABG</td>
<td>105 MiECC /139 CCPB (high-risk patients)</td>
<td>Type I CK levels were significantly lower 6 h after surgery in the MiECC group ($P &lt; 0.05$). Need of red blood cell transfusion was significantly lower after MiECC surgery ($P &lt; 0.05$). 30-day mortality was significantly lower in the MiECC group ($P &lt; 0.01$).</td>
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<tr>
<td>Kolat et al., J Cardiothorac Surg, 2014 [50]</td>
<td>Retrospective cohort analysis</td>
<td>CABG</td>
<td>1137 MiECC /1137 CCPB</td>
<td>Type I Postoperative requirement of renal replacement therapy ($P = 0.01$), respiratory insufficiency ($P = 0.004$) and incidence of low cardiac output syndrome ($P = 0.003$) were significantly increased in patients with CCPB. 30-day mortality was reduced in patients with MiECC ($P = 0.03$). ICU stay ($P = 0.70$), hospital stay ($P = 0.40$) and postoperative low cardiac output syndrome ($P = 0.83$) did not show significant differences between both groups.</td>
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<tr>
<td>Ried et al., J Cardiothorac Surg, 2013 [51]</td>
<td>Propensity score analysis</td>
<td>Emergency CABG</td>
<td>146 MiECC /175 CCPB</td>
<td>Type I Stroke rate was significantly higher among CCPB patients ($P = 0.026$). In-hospital mortality, combined adverse end-point rate, postoperative bleeding and need for transfusion were statistically insignificant in the study groups.</td>
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<tr>
<td>Koivisto et al., Perfusion, 2010 [52]</td>
<td>Propensity score analysis</td>
<td>CABG</td>
<td>89 MiECC /147 CCPB</td>
<td>Type II 30-day mortality was reduced in patients with MiECC ($P = 0.03$). ICU and hospital stay were comparable between the two groups.</td>
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<tr>
<td>Anastasiadis et al., Int J Cardiol, 2013 [53]</td>
<td>Cost-analysis</td>
<td>CABG</td>
<td>1026 MiECC/1023 CCPB</td>
<td>Type II Focus on cost-effectiveness</td>
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<tr>
<td>Fernandes, Perfusion, 2010 [54]</td>
<td>Retrospective cohort study</td>
<td>CABG</td>
<td>15 MiECC</td>
<td>Type II Using lower than predicted flows, adequate perfusion was provided.</td>
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<tr>
<td>Puehler et al., Thorac Cardiovasc Surg, 2010 [55]</td>
<td>Retrospective comparative cohort study</td>
<td>CABG</td>
<td>119 MiECC /119 CCPB</td>
<td>Type I MiECC patients had a tendency towards a lower 30-day mortality rate, a better postoperative renal function and reduced ventilation times. CPB time and postoperative high-dose inotropic support were significantly lower in the MiECC group. ICU and hospital stay were comparable between the two groups.</td>
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</table>

- **a-GST:** a-glutathione S-Transferase
- **AF:** atrial fibrillation
- **AKI:** acute kidney injury
- **AVR:** aortic valve replacement
- **CABG:** coronary artery bypass grafting
- **CCPB:** conventional cardiopulmonary bypass
- **CPB:** cardiopulmonary bypass
- **HFABP:** heart type fatty acid binding protein
- **ICU:** intensive care unit
- **IFABP:** intestinal type fatty acid binding protein
- **IL:** interleukin
- **MAP:** mean arterial pressure
- **MiECC:** minimal invasive extracorporeal circulation
- **MVR:** mitral valve replacement
- **NGAL:** neutrophil gelatinase-associated lipocalin
- **OPCAB:** off-pump coronary artery bypass grafting
- **POD:** postoperative day
- **RCT:** randomized controlled trial
- **SIRS:** systemic inflammatory response syndrome
- **TNF:** tumour necrosis factor
- **TnT:** troponin-T
- **TnI:** troponin I
circulation (mECC), minimized extracorporeal circulation, mini-cardiopulmonary bypass (mCPB, mini-CPB), minimally invasive cardiopulmonary bypass (MICPB), miniaturized cardiopulmonary bypass (MCPB), veno-arterial extracorporeal membrane oxygenation, minimized perfusion circuit, minimized extracorporeal life support system, minimized CPB, MiECC. This divergent terminology creates confusion and disagreement between centres. But the major problem is the fact that the focus is made only on the priming volume of the circuit and not on the reduction of the adverse effects of ECC.

The Steering Committee of MiECTiS uses the term 'minimal invasive' to describe a procedure which involves not only the CPB circuit, but the global approach to the procedure. This concept strives to render the procedure minimally invasive as opposed to the widely employed misnomer 'minimal invasive' when a limited surgical access is performed. The term ‘minimal invasive’ is misleading since the patient is often a longer period on CPB, cross-clamping and duration of the anaesthesia are prolonged. In this sense, the term ‘minimal invasive extracorporeal circulation’ corresponds better to the above mentioned concept and should be used to describe this technology with the abbreviation: ‘MiECC’.

### Components of MiECC system

In order to be characterized as MiECC, the main components of the system must include a closed CPB circuit; biologically inert blood contact surfaces; reduced priming volume; a centrifugal pump; a membrane oxygenator; a heat exchanger; a cardioplegia system; a venous bubble trap/venous air removing device and a shed blood management system.

Because different groups have utilized either commercially available or customized CPB circuits with a variety of components, the Consensus Meeting defined the main components of the CPB circuit when it should correspond to a MiECC system. The Steering Committee of MiECTiS emphasizes that a MiECC system should comprise all necessary elements to obtain a maximal benefit.

Originally, MiECC system was an extracorporeal life support (ECLS) circuit with the possibility to administrate cardioplegia (Type I) and used mainly to perform CABG procedures [10]. However, the safety concerns regarding air entrapment/air lock into the venous line prompted the integration of venous bubble trap/venous air removing devices into the system (Type II). This design increased safety for CABG procedures and enabled aortic valve surgery [11].

### Table 3: Summary of statements endorsed by the Expert Committee

<table>
<thead>
<tr>
<th>Recommendation</th>
<th>Level of evidence</th>
<th>References</th>
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<tbody>
<tr>
<td>MiECC refers to a combined strategy of surgical approach, anaesthesiological and perfusion management and should not be limited to the CPB circuit alone. In order to be characterized as MiECC, the main components of the system must include closed circuit; biologically inert blood contact surfaces; reduced priming volume; centrifugal pump; membrane oxygenator; heat exchanger; cardioplegia system; venous bubble trap/venous air removing device; shed blood management system. Additional components that can be integrated to a MiECC system are pulmonary artery vent; pulmonary vein vent; aortic root vent; soft bag/soft-shell reservoir; hard-shell reservoir (modular systems); regulated smart suction device; arterial line filtration.</td>
<td><strong>A</strong></td>
<td>[17, 19, 20]</td>
</tr>
<tr>
<td>MiECC systems reduce haemodilution and better preserve haematocrit as well as reduce postoperative bleeding and the need for RBC transfusion</td>
<td><strong>A</strong></td>
<td>[17, 19, 20]</td>
</tr>
<tr>
<td>MiECC systems reduce the incidence of postoperative atrial fibrillation</td>
<td><strong>A</strong></td>
<td>[17, 19, 20]</td>
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<tr>
<td>MiECC systems preserve renal function</td>
<td><strong>A</strong></td>
<td>[20, 39]</td>
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<tr>
<td>MiECC is associated with improved myocardial protection</td>
<td><strong>A</strong></td>
<td>[20, 43-45]</td>
</tr>
<tr>
<td>Class IIA: Inflammatory response assessed by specific inflammatory markers is attenuated with use of MiECC</td>
<td><strong>B</strong></td>
<td>[23-26]</td>
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<tr>
<td>MiECC systems can reduce cerebral gaseous microembolism and preserve neurocognitive function</td>
<td><strong>B</strong></td>
<td>[29, 30]</td>
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<td>MiECC exerts a subclinical protective effect on end-organ function (lung, liver, intestine) which is related to enhanced recovery of microvascular organ perfusion</td>
<td><strong>B</strong></td>
<td>[39, 46-48]</td>
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<td>Class IIB: Within a MiECC strategy, less thrombin generation may permit reduced heparin dose targeted to shorter ACT times. When such a strategy is followed, individual heparin dose should be determined using heparin dose-response monitoring systems</td>
<td><strong>B</strong></td>
<td>[14, 15, 56, 57]</td>
</tr>
<tr>
<td>MiECC appears to offer survival benefit in terms of lower 30-day mortality after CABG procedures</td>
<td><strong>B</strong></td>
<td>[20, 49, 50, 51]</td>
</tr>
<tr>
<td>The use of short-acting opioids in combination with propofol or volatile anaesthetics, and hypnotic effect monitoring by processed EEG, is recommended for induction and maintenance of anaesthesia for MiECC-based surgery. TOE findings pertinent to institutional management of MiECC should be communicated during the preoperative surgical safety time out</td>
<td><strong>B</strong></td>
<td>[16, 58-61]</td>
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ACT: activated clotting time; CABG: coronary artery bypass grafting; EEG: electroencephalogram; MiECC: minimal invasive extracorporeal circulation; RBC: red blood cells; TOE: transthoracic echocardiography; rSo2: regional cerebral oxygen saturation.
The need for blood volume management during valvular procedures required the addition of a soft-bag/soft-shell reservoir integrated into the system (Type III). This enabled safe performance of aortic valve surgery and other intracardiac procedures. Initiation of modular MiECC (hybrid) systems that integrate a second open circuit with a venous reservoir and cardiomyotomy suction as a stand-by component (Type IV) enabled performance of complex procedures that pertain a high possibility of unexpected perfusion scenario [12, 13]. Classification of MiECC types is illustrated in Fig. 1. The Consensus Meeting defined as a prerequisite for a system to be considered as MiECC to have at least Type II circuit characteristics.

Additional components to be integrated into a MiECC system are (i) pulmonary artery vent, (ii) aortic root vent, (iii) pulmonary vein vent, (iv) soft-bag/soft-shell reservoir, (v) hard-shell reservoir (modular systems), (vi) regulated smart suction device and (vii) arterial line filtration.

Modular systems

The major reticence to limit expansion of MiECC is due to the thoughts about safety in case of massive air entrance into the system or significant blood loss. Although CABG and valve surgery are feasible with the standard type II MiECC circuit, a modular configuration is welcome to expand MiECC for the majority of cardiac procedures and to create a 'safety net' for unexpected intraoperative scenarios. Recently published results from a single-centre indicate that a modular circuit design offers 100% technical success rate in high-risk patients, even in those undergoing complex procedures including reoperations, valve and aortic surgery as well as emergency cases [12].

Anticoagulation management

During perfusion with MiECC, less thrombin generation may allow reduced heparin dose targeted by shorter ACT (Class of Recommendation IIB, Level of Evidence B). In this case, individual heparin dosage should be determined using heparin dose-response monitoring systems.

A number of factors including better biocompatible surfaces, elimination of blood-air interaction and exclusion of unprocessed shed blood reinfusion favourably influence thrombin generation under MiECC system compared with the standard CPB [56]. A patient-adjusted and/or a procedure-adjusted coagulation management based on unfractionated heparin (UFH) can be adopted.

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**Figure 1:** Classification of MiECC circuits [12]. [Note that the modular type IV circuit is literally type III with a standing-by component, used only when necessary] (X: pump; O: oxygenator; C: cardioplegia; T: bubble trap/air-removing device; V: vent (aortic/pulmonary); S: soft-bag/reservoir; H: hard-shell/reservoir). MiECC: minimal invasive extracorporeal circulation.
Anaesthesia for surgery on MiECC

The use of short-acting opioids in combination with propofol or volatile anaesthetics, and monitoring of the depth of anaesthesia by processed electroencephalogram (EEG), is recommended if all patients undergoing cardiac surgery with MiECC (Class of Recommendation IIB, Level of Evidence C). Transoesophageal echocardiography (TOE) findings pertinent to institutional management of MiECC should be communicated during the preoperative surgical safety time out (Class of Recommendation IIB, Level of Evidence C).

Anaesthetic management of patients undergoing cardiac surgery with the aid of a MiECC system follows the international recommendations, especially regarding the use of TOE [58, 59]. Following anaesthesia induction, TOE may provide additional information that may influence the site and/or the type of cannulation or perfusion strategy (e.g., patent foramen ovale, significant mitral or aortic valve pathology or severe aortic atheromatosis). This information is important when Type I or II MiECC systems are used, whereas any modifications can be accommodated when Type III or modular type IV configuration are available.

Specifically, the absence of venous reservoir in MiECC systems renders the patient’s own venous capacitance compartment critical for haemodynamics as well as for optimal volume management. Positioning of the patient (Trendelenburg or anti-Trendelenburg) and low-dose vasoactive agents are useful to control intraoperative haemodynamics. Excessive fluid administration should be avoided to reduce haemodilution and avoid transfusion [63].

Beneficial effects of MiECC include attenuation of inflammatory response, higher haematocrit, less coagulation disorders and improved end-organ function (brain, kidneys, lungs). It facilitates implementation of fast-track protocols [16]. Hence, perioperative use of short-acting intravenous and/or volatile anaesthetic agents is recommended. Moreover, the titration of anaesthetic agents using processed EEG ensures adequate anaesthesia depth [60]. Microporous capillary membrane oxygenators enable volatile anaesthetics to be used for anaesthesia maintenance, which is not feasible with diffusion membrane oxygenators [61]. To date, RCTs comparing different anaesthetic protocols for MiECC-based surgery are still missing.

Haemodilution: haematocrit—transfusion

MiECC systems reduce haemodilution, better preserve haematocrit and reduce postoperative bleeding and the need for RBC transfusion (Class of Recommendation I, Level of Evidence A).

There is compelling evidence that MiECC—mainly because of the significantly reduced priming volume of the circuit—reduces haemodilution and results in a higher haematocrit at the end of the perfusion period [17, 18]. This significantly reduces need for red blood cells transfusion and improves oxygen delivery during perfusion [13, 17, 19, 20, 65]. Coagulation disorders are reduced [17] and platelet count and function are better preserved following perfusion with MiECC systems [21]. Postoperative bleeding and incidence of re-exploration are significantly lower in patients operated with MiECC [20]. As it reduces haemodilution, MiECC fulfils Class of Recommendation I, Level of Evidence A indication for blood conservation according to the STS guidelines, especially in patients at high risk for adverse effects of haemodilution (paediatric patients and small-sized adults) [8]. Patients refusing transfusion of allogeneic blood products, e.g. Jehovah’s Witnesses, are optimal candidates for this strategy [22].

Attenuation of the inflammatory response

Inflammatory response is attenuated with use of MiECC (Class of Recommendation IIA, Level of Evidence B).

Several studies have investigated the inflammatory response triggered conventional CPB and compared it with MiECC systems. MiECC components are designed to limit the severity of SIRS. Coating and reduction of the size of the circuit reduce the amount of foreign surfaces, which is the main trigger of SIRS. Other studies still have to confirm this observation [66]. Assessment of the inflammatory response is complex and clinical presentation is highly variable [67]. Nevertheless, some studies provide evidence of the beneficial effects of MiECC. Moreover, Fromes et al. [23] described a less pronounced intraoperative decrease of monocytes as well as during the first 24 h in patients with MiECC than in those with conventional CPB. Others demonstrated significantly lower peak levels of IL-6 under MiECC [23, 24, 68]. Finally, several studies demonstrated that perfusion with MiECC resulted in significantly lower levels of neutrophil elastase—a specific marker of neutrophil activation—than with conventional CPB [23, 25, 26].

Neurological function

MiECC systems reduce cerebral gaseous microembolism and better preserve neurocognitive function (Class of Recommendation IIA, Level of Evidence B).

Several prospective studies and meta-analyses have reported reduced incidence of stroke following MiECC when compared with conventional CPB [19, 27, 28]. A recent meta-analysis found a trend to reduction of neurological damage in favour of MiECC [20]. Of course, stroke is multifactorial and the perfusion system is only one of the issues besides aortic manipulations and other patient’s specific factors [69]. A possible explanation for the neuroprotective effect of MiECC is the significant reduction of gaseous microemboli [29–33]. MiECC also offers improved cerebral perfusion during CPB, as indicated by the lower reduction in near infrared spectroscopy (NIRS)-derived regional cerebral oxygen saturation values and cerebral desaturation episodes [29, 32, 34, 35]. Reduced incidence of cerebral desaturation episodes favourably affects neurocognitive outcome [70–72].

Atrial fibrillation

MiECC reduces the incidence of postoperative atrial fibrillation (AF) (Class of Recommendation I, Level of Evidence A).
Several randomized studies have demonstrated that postoperative AF is significantly reduced following MiECC when compared with conventional CPB [13, 16, 24, 36]. Moreover, there is strong evidence of a lower incidence of AF in all meta-analyses regarding MiECC systems [19, 20, 28]. Attenuated inflammatory reaction and less volume shifts associated with MiECC may be an explanation for this beneficial effect [37].

Renal function

MiECC preserves renal function (Class of Recommendation I, Level of Evidence A).

Several studies have shown that the use of MiECC systems was associated with better preservation of renal function [38–40]. This was confirmed by a meta-analysis of 24 RCTs but this meta-analysis and other studies failed to demonstrate a reduced incidence of postoperative renal failure [20, 38, 41]. More stable haemodynamics together with higher perfusion pressure and a reduced need for vasopressors during MiECC perfusion may explain this observation [10, 42]. A significant independent association was found between the lowest haematocrit value during bypass and acute renal injury, with significant benefits on renal function seen after reduction of the priming volume. This may be due to a higher DO2 associated with a higher haematocrit on CPB [65]. In addition, different markers to evaluate renal function (i.e. glomerular filtration rate, levels of neutrophil gelatinase-associated lipocalin) confirm better renal protection under MiECC. Larger studies are required to investigate if this protective effect is sufficient to prevent development of acute renal failure.

Myocardial protection

MiECC is associated with improved myocardial protection (Class of Recommendation I, Level of Evidence A).

Several studies have demonstrated a beneficial effect of MiECC on intraoperative myocardial protection [10, 20, 43, 44]. Reduced cardioplegia volumes with less crystalloids and attenuation of SIRS may explain this beneficial effect [23]. Studies with MiECC and intermittent cross-clamping show a similar effect on myocardial protection [45]. However, myocardial protection is not related only to the duration of ischaemia, but also to the reperfusion phase. Increased arterial pressure during CPB as well as the volume-constant perfusion with a closed system may also contribute to improved myocardial protection [38, 42].

End-organ protection

MiECC has a subclinical protective effect on end-organ function (lung, liver, intestine) caused by improved microvascular organ perfusion (Class of Recommendation IIa, Level of Evidence B).

MiECC is a closed system that allows a better peripheral perfusion with higher arterial pressure and systemic vascular resistance close to normal values [38]. This is associated with reduced requirement for vasoactive support [10, 42]. Data from randomized studies suggest improved lung protection [46], attenuated liver and intestinal dysfunction [39, 46, 47]. These studies evaluated only surrogate markers of end-organ dysfunction that may benefit from MiECC, whereas the effects remain subclinical. However, it may become clinically perceptible in high-risk patients and in those with longer procedures since MiECC would lead to fewer alterations of microperfusion [48].

Mortality

MiECC appears to offer survival benefit in terms of lower 30-day mortality after CABG procedures (Class of Recommendation IIB, Level of Evidence B).

A number of studies have demonstrated a trend towards reduced mortality in CABG performed on MiECC. A recent meta-analysis of 24 studies involving 2770 patients showed that MiECC was associated with a significant decrease in mortality, compared with conventional CPB (0.5 vs 1.7%; P = 0.02) [20]. This finding has also confirmed by other studies [49, 50, 51]. A trend towards decreased mortality in favour of MiECC has also been found in meta-analyses [19, 28] and in a propensity score analysis [52]. This survival benefit may be the result of the cumulative beneficial effects of MiECC on end-organ protection but it calls for a multicentre RCT sufficiently powered to prospectively investigate this survival benefit.

Cost-effectiveness

Data from a cost-analysis study indicate a cost-effectiveness of MiECC systems that offer economic advantages in various healthcare settings [53]. Nevertheless, these results have to be considered in the context of the local conditions. A more detailed analysis together with an analysis from a payer’s perspective is necessary. Better standardization should be achieved to allow comparison of costs and economic benefits.

DISCUSSION

MiECC systems have been developed to integrate all advances in CPB technology in one closed circuit: the goal is to improve biocompatibility and minimize side-effects of CPB. MiECC is associated with more stable haemodynamics during and early after perfusion and better end-organ protection. This concept provides comparable or better outcomes in terms of morbidity and mortality in CABG and valve procedures, as shown in prospective randomized studies and meta-analyses. However, despite several clinical advantages, the implementation of MiECC technology remains weak probably there are still some concerns regarding air handling as well as blood and volume management during perfusion [12]. This Consensus paper primarily serves to summarize the available information about this technology and to clarify some of the open issues. We have made substantial efforts to provide the best available actual evidence and strongly encourage readers to consider the technology as a multidisciplinary strategy.

There is still debate about the optimal handling of air during the perfusion, as well as volume and blood management when a MiECC system is used. Mean arterial pressure (MAP) is usually higher during MiECC: this raises the question of optimal pump flow rate during MiECC perfusion [10, 42]. A reference blood flow based on body surface area is not a guarantee of adequate body perfusion during CPB. Modern protocols adjust pump flow to achieve adequate DO2. In this area, it is still unclear if the use of MiECC may allow lower than traditional cardiac index without end-organ damage as has been suggested by recent studies [54, [54]
The use of NIRS and other parameters to monitor cerebral blood flow may lead to greater individualization of perfusion index for adequate end-organ perfusion [35, 74]. Lower heparin requirement and reduced haemodilution offered by MiECC facilitate the management of postoperative bleeding. The prophylactic use of low-dose antifibrinolytics [75] and POC coagulation management based on thromboelastometry and aggregometry is generally advised [76]. In patients with higher perioperative risk [52], those with low ejection fraction and emergencies [51, 52, 55], MiECC has proved to be safe.

In general, MiECC can be considered as the ‘circuit-of-choice’ to replace conventional CPB at least for CABG surgery. Novel modular systems (Type IV MiECC) may be utilized for all cardiac procedures. We believe that the terms ‘circuit’ which refers to the CPB, the ‘MiECC system’ which integrates certain components to a CPB circuit and the ‘MiECC strategy’ that represents the multidisciplinary approach to MiECC should be differentiated. The MiECTiS advocates this strategy to obtain the maximal benefits for the patients. The authors believe that MiECC should be understood as an additional tool in the chapter of minimal invasiveness. The latter should not be restricted to ‘minimal-access’ surgery, but should also incorporate a strategy towards a ‘more physiologic CPB’. The use of MiECC should be integrated within fast-track algorithms, POC management of coagulation disorders together with any initiatives that improve aortic assessment (epiaortic ultrasound), novel anti-inflammatory strategies, low shear-stress cannula design and implementation of contemporary biofiltration techniques.

Lack of high-volume data requires the creation of a registry to further evaluate this technology. Moreover, the variation in extent of miniaturization/complexity of MiECC systems should be analysed. Additional RCTs, focusing on valve and other cardiac procedures, as well as large cohorts of patients will provide more evidence regarding clinical effectiveness. Adequately powered prospective multicentre studies are required in order to prove the superiority of the MiECC over the conventional CPB.

Concerns in the literature have been raised regarding decreased safety, ventricular dilatation during perfusion using the MiECC circuit, loss of a bloodless field and the risk of air embolism [77, 78]; however, these reports are anecdotal and are not supported by large-scale studies. Loss of safety during perfusion with a modern MiECC circuit is easily addressed with integration of a venous bubble trap/air removing device into the circuit. Moreover, significant air entrainment that blocks the circuit could be resolved immediately by a skilled perfusionist. Ventricular dilatation, attributed to poor off-loading of the heart, is anticipated with the use of aortic root and/or pulmonary artery/vein venting from Type II MiECC onwards. The same applies to creation of a full bloodless field. Special patient populations, such as patients with a higher body surface area requiring higher circulatory flows, are easily managed with kinetic-assisted venous drainage and increased flow through the centrifugal pump. Regarding air embolism, contemporary evidence suggests that there is significantly reduced amount of gaseous microemboli in the arterial line of MiECC systems compared with conventional CPB [79].

Nevertheless, it should be emphasized that MiECC is a demanding system that should be implemented in cardiac surgery as a strategy and not as a simple circuit. Real teamwork from all disciplines of the surgical team, meticulous surgery, a skilful perfusionist and optimal anaesthetic management are mandatory for a more physiological perfusion that could lead to improved clinical outcomes. MiECTiS supports initiatives that promote research and clinical application of MiECC systems as a strategy through multidisciplinary training programmes (dry labs/hands-on simulators, wet labs, peer-to-peer workshops). Integration of specific training programmes under the accreditation of MiECTiS will stimulate and improve the collaboration between clinicians while the industry will get important information to further improve the systems. MiECTiS is planning to endorse a comprehensive and structured programme that contributes to the advancement of patient care.

In conclusion, the authors consider MiECC as a physiologically based strategy and not just a CPB circuit or a particular product. For this reason, multidisciplinary approach is mandatory. Collaboration between surgeons, anaesthesiologists and perfusionists is of paramount importance to emphasize the key tenets of MiECTiS.

SUPPLEMENTARY MATERIAL

Supplementary material is available at ICVTS online.

REFERENCES


