Dear Sir,

Although kidney transplantation has become an effective means of treating end-stage renal disease (ESRD), many patients die waiting for organ transplant. This problem is not an isolated phenomenon in Poland (13.6 donors per 1 million inhabitants) but in other European countries, despite the much greater number of organs transplanted each year (from 23.4 donors per 1 million inhabitants in the Czech Republic 40.2 per million in Spain), Poland continues to seek and create opportunities for the development program of organ procurement and transplantation. The alternative option is the inclusion donor after circulatory death (DCD) in the transplantation program, for which circulatory criteria are used to define death. The complicated procedure required to maintain the viability of organs for transplantation and dramatic time criteria in the absence of appropriate organizational capabilities have caused that transplantation from DCD donors is not yet widespread in Poland [1–3].

We present the procedural preparation for the Poland’s first case of organ (kidney) transplantation from a circulatory death donor in which perfusion was supported by Extracorporeal Membrane Oxygenation (ECMO). The procedure is a component of the “ECMO for Wielkopolska” program established to serve 3.5 million inhabitants of the Wielkopolska region in Poland. Because this organizational model is complex and expensive, we used advanced

FIGURE 1. SCA in public-ALS on MCC

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high-fidelity medical simulation testing in order to prepare for real-life experience (Fig. 1, 2) [4–6].

The system of self-made silicone loops and tubing simulating blood vessels (modified polyethylene), filled with pressurized red-dyed liquid was implanted into the groin of a mannequin and covered by artificial skin. The real time scenario included all the following crucial steps: prehospital identification; CPR ALS; perfusion therapy (CPR-ECMO or DCD-ECMO); inclusion and exclusion criteria matching; suitability for mechanical chest compression AUTOPULS (ZOLL, Chelmsford, USA); DCD confirmation and donor authorization (Fig. 3–5). The warm ischemia time, i.e. time from the first contact of mannequin to the cannulation of artificial vessels and starting in situ organ perfusion using CARDIOHELP (Maquet, Rastatt, Germany), including CPR, did not exceed two hours. In our case of a DCD-ECMO procedure, performed for the first time in Poland, its success is reassuring and helps to fight obstacles and establish a functioning programme. Soon after a successful advanced simulation test, four DCD-ECMO procedures I Maastricht category II (unsuccessful resuscitation) were performed resulting in two double successful kidney transplantations for four recipients.

In conclusion, during debriefing, it was found that the previous training allowed one to build a procedural chain, eliminate errors at the stage of identification, notification, transportation, qualifications and ECMO organs perfusion. The training program resulted in a team appropriately qualified to successfully undertake this complex procedure. Moreover, one part of the “ECMO for Wielkopolska” program is the learning website: www.ecmo.pl.

**Conflict of interest:** None declared.

**REFERENCES**


