Altered pharmacological effects of adrenergic agonists during hypothermia

Erik Sveberg Dietrichs¹,²*, Georg Sager³ and Torkjel Tveita¹,⁴

Abstract

Rewarming from accidental hypothermia is often complicated by hypothermia-induced cardiac dysfunction, calling for immediate pharmacologic intervention. Studies show that although cardiac pharmacologic support is applied when rewarming these patients, a lack of updated treatment recommendations exist. Mainly due to lack of clinical and experimental data, neither of the international guidelines includes information about pharmacologic cardiac support at temperatures below 30 °C. However, core temperature of accidental hypothermia patients is often reduced below 30 °C. Few human studies exploring effects of adrenergic drugs during hypothermia have been published, and therefore prevailing information is collected from pre-clinical studies. The most prominent finding in these studies is an apparent depressive effect of adrenaline on cardiac function when used in doses which elevate cardiac output during normothermia. Also noradrenaline and isoprenaline largely lacked positive cardiac effects during hypothermia, while dopamine is a more promising drug for supporting cardiac function during rewarming. Data and information from these studies are in support of the prevailing notion; not to use adrenergic drugs at core temperatures below 30 °C.

Keywords: Hypothermia, Cooling, Rewarming, Rearming shock, Pharmacology, Cardiovascular dysfunction, Adrenergic drugs, Inotropic, Vasopressor

Background

Rewarming victims of accidental hypothermia is often complicated by hypothermia-induced cardiac dysfunction. In its fulminant form this condition is described as rewarming shock; an acute heart failure with a progressive fall in cardiac output (CO) where the patient terminates in a sudden and intractable fall in blood pressure [1]. This serious complication to clinical therapy adds to the virtually unchanged low survival rate of accidental hypothermia over the last decades [2, 3]. Cardiac supportive therapy has to be instituted during rewarming in an attempt to prevent the imminent cardiovascular collapse. Inotropic drugs, i.e. drugs that enhance force of cardiac contraction [4] could provide such pharmacologic support, but current guidelines do not support this view. Both the American Heart Association and the European Resuscitation Council advise against using drugs like adrenaline below 30 °C [5, 6]. Studies investigating patient treatment do however report that inotropic drugs are administered during rewarming in 47–66% of patients [7, 8]. Only about 10% of patients with acute heart failure caused by disease or events other than hypothermia receive the same treatment [9]. This lack of consensus-based guidelines on cardiac inotropic support in hypothermic patients cause confusion, even within the British health care system [10]. Importance of finding optimal treatment for these patients is manifest from that survival is possible after extreme exposure to hypothermia. Case-reports show that early resuscitation enables survival after cooling to 13.7 °C [11] or cardiac arrest close to 7 h [12], but mortality rate is still reported at 30% [2]. Seeking evidence-based pharmacologic treatment options, we have explored the literature for pre-clinical and clinical studies on use of inotropic drugs during hypothermia and rewarming, with interesting findings. Most studies, including several from our group, focus on adrenergic receptor agonists. The information provided, presents a valuable insight in hypothermia-induced changes in cardiovascular pharmacology, laying...
Molecular effects

The effect of phosphorylation of cardiac troponin I enhances myocardial calcium influx, thus enabling contraction, while phosphorylation of sarcolemmal L-type calcium channels gives increased calcium increase. This activates protein kinase A (PKA), stimulating of adenylyl cyclase, which elevates cyclic AMP (cAMP). This activates protein kinase A (PKA), which phosphorylates several proteins.

Methods

The aim of this paper was to describe effects of adrenergic agonists during hypothermia. Relevant publications were found through literature search, using PubMed (Medline) and Google Scholar search engines. Experimental and clinical studies included in this narrative review were selected according to their relevance by the authors, who all have a special interest in hypothermia and pharmacology. Reference lists of included papers were studied to discover publications that were not detected through use of search engines. Additional articles describing pathophysiology of hypothermia, treatment guidelines and general knowledge on adrenergic agonists and receptor function, were included for background information.

Adrenergic receptors (Table 1)

Extracellular binding of adrenergic receptor agonists facilitates initiation of intracellular processes through G-protein coupled signalling. In 1948, Ahlquist described how the adrenergic receptors are divided into two main groups [13], namely α- and β-receptors. Subgroups have later been identified and separated the receptors into α1, 2 and β1–3p, which have a broad variety of effects, among them hemodynamic. The β1-receptor is considered most important for inotropic effect and is also more numerous in the mammalian heart (75%) than β2 and β3 [14]. β1-stimulation enhances heart muscle contraction, increase heart rate and enhance relaxation of myocardial tissue [15]. The effect of β1-agonists is mediated through stimulation of adenylyl cyclase, which elevates cyclic AMP (cAMP). This activates protein kinase A (PKA), which phosphorylates several proteins. Phosphorylation of sarcolemmal L-type calcium channels gives increased calcium influx, thus enabling contraction, while phosphorylation of cardiac troponin I enhances myocardial relaxation [14]. β2-stimulation most importantly gives vasodilation due to smooth muscle relaxation [16]. Both α1-receptors are divided into three subgroups. Stimulation of all α1-receptors will in general induce smooth muscle contraction, which gives vasoconstriction [17]. The α2-receptor subgroups have differing abilities, among them both vasodilation and vasoconstriction [18]. Cardiovascular effects of adrenergic stimulation are also transmitted through dopamine receptors by direct effect on smooth muscle, giving vasodilation mediated by D1-like receptors and indirectly by D2-like receptors. Several dopamine receptor subtypes are located in the human heart, namely D1, D2, D4 and D5 [19], but stimulation of these receptors does not have a pronounced effect on cardiac contractility in rats [20]. Adrenergic receptor agonists have varying affinity for β1, α1 and dopamine-receptors and their subgroups, explaining their distinct properties.

β1-receptor function during hypothermia

As the β1-receptor is considered most important for providing inotropic effect in normothermic conditions [14] studies on administering adrenergic drugs to ameliorate rewarming shock have also targeted this receptor. In a study from our lab we have reported a 4-fold increase of in vivo cardiac cAMP content during β1-receptor stimulation at 15 °C, showing that β1-receptor function is not depressed at low temperatures. This was confirmed by an in vitro 9-fold increase of β1-receptor sensitivity in isolated rat cardiomyocytes cooled to 15 °C [21]. Such hypothermia-induced in vitro β1-receptor super-sensitivity was also described in isolated heart preparations from guinea pig [22, 23] and rabbit hearts at 22 °C [24]. Results from intact (in vivo) animal experiments studying effects of adrenergic receptor-ligands and -blockers indicate that both β1- and β2-receptor function is maintained during cooling to 28–30 °C, but that response to agonist binding is depressed by cooling below 30 °C [25–27]. The relationship between an apparent increase in β-receptor sensitivity and decreased inotropic effect below a core temperature

| Table 1 Adrenergic receptors |
|-----------------------------|
| α-receptor  | β-receptor | Dopamine-receptor |
| Subgroups       | α1, α2    | β1, β2, β3     | D1, D2, D3, D4, D5 |
| Mechanism of action | G-protein coupled receptors | G-protein coupled receptors | G-protein coupled receptors |
| Molecular effects | α1: Activation of PLC, IP3 mediated intracellular calcium increase | Increase of cAMP, protein phosphorylation (β1, β2, β3) intracellular calcium increase (β1) or decrease (β2) | D1, D2: Increase of cAMP D2: Decreased cAMP production |
| Hemodynamic effect | α1: Vasoconstriction† | β1: Heart rate†, contractility† | D1, D2: Vasodilation† |
| Dominant location | α1: Smooth muscle | β1: Cardiac tissue | D1, D2, D3, D4, D5 |
|                          | α2: Central nervous system | β2: Smooth muscle | Central nervous system |
of 30 °C is uncertain. Mann [28] found excessive cAMP levels, seen during β1-receptor stimulation in hypothermic rats [21], to be cardiotoxic, through initiating unphysiological increase of cytosolic calcium levels, mediated by increased phosphorylation of L-type calcium channels [28]. Hypothermia-induced calcium overload is also known to take place in response to prolonged hypothermia in rats per se [29, 30] and an additional increased calcium load in response to a pharmacologic stimulation might explain lacking effect of β1-agonists during hypothermia. Another observed effect of temperature reduction in rats is reduced myocardial calcium sensitivity due to hypothermia-induced elevation of PKA-mediated phosphorylation of Ser 23/24 at cardiac troponin I [31]. Increased levels of cAMP will increase such PKA-mediated Ser 23/24 phosphorylation and give a negative inotropic effect. Such phosphorylation will also enhance cardiac relaxation [32], which is normalized during rewarming from hypothermia as diastolic function is restored [33]. Thus, favorable or harmful effects of β1-agonists administration in hypothermic subjects appears associated to inotropic rather than lusitropic properties, as hypothermia-induced cardiac dysfunction is an isolated impairment of systolic function [1].

The aforementioned studies do however show differences in species and experimental conditions and use several β1-receptor agonists with varying properties, including α-receptor agonism that promotes vasoconstriction. A resulting increase of systemic vascular resistance (SVR) cause a pronounced negative effect on cardiac function in rats during hypothermia [21, 27, 34, 35]. For assessment of clinical properties of these drugs, it is therefore necessary to evaluate their individual pharmacologic effects when used during hypothermia.

Adrenergic receptor agonists (Table 2, Fig. 1)

Adrenaline
Adrenaline will enhance cardiac contraction and heart rate and either decrease (low-dose) or increase (high-dose) SVR in normothermic conditions [36], conducted by loss of β-adrenergic selectivity at low doses with increasing α-stimulation at higher doses.

Adrenaline during hypothermia Pharmacodynamic effects of adrenaline do not seem to be independent of temperature changes. Rubinstein found that doses inducing vasodilation in normothermic dogs would give increased SVR during hypothermia and stated that the inotropic effect of adrenaline is reduced at 25 °C [37]. A similar study on rats conducted in our lab, showed that a high dose of adrenaline (1.25 μg/min) increased stroke volume (SV) and CO in normothermic animals. When an equal dose was administered during rewarming from 15 °C however, SV and CO were unaffected. In contrast, a low dose (0.125 μg/min) adrenaline, which induced vasodilation during normothermic conditions, failed to reduce SVR or mean arterial pressure (MAP) during rewarming, but led to an elevated CO [34]. The positive effect of low-dose adrenaline during hypothermia has also been reported from experiments using dogs [38]. We found the same dose–response relationship during cooling, where 0.125 μg/min but not 1.25 μg/min adrenaline gave positive cardiac effects during cooling to 28 °C. After rewarming, only rats that had received saline during cooling showed pre-hypothermic hemodynamic responses to adrenaline [27]. An additional study on rats from our lab showed that 1 μg/min of adrenaline given during cooling caused a maintained depression of cardiac function during rewarming [35]. These results indicate that hypothermia has a severe impact on cardiac inotropic effects mediated by the β1-receptor pathway, as β1-adrenergic stimulation during hypothermia also has a negative impact on inotropic effect of β1-agonists after rewarming. The same phenomenon is also observed in a feline model of hypothermia and rewarming [39, 40]. From a combined in vitro and in vivo study in our lab, we showed that this hypothermia-induced reduction of inotropic effect via β1-receptor stimulation is seen in the presence of in vivo and in vitro β1-receptor super-sensitivity. Both increased β1-receptor binding and elevated cAMP levels were seen during administration of β1-receptor ligands in hypothermic conditions, as compared to normothermia. This study indicates that the detrimental effects of adrenaline during hypothermia is a consequence of adrenaline-induced increase in SVR via peripheral vascular α-receptor stimulation [21]. Failure of low doses of adrenaline to reduce MAP during hypothermia [34], further implies the presence of increased α-receptor agonism, or reduced effect of β2-receptor agonism in hypothermia.

Altogether, the present hypothermia-induced alteration of pharmacodynamic effects and an expected narrowed therapeutic window of adrenaline, advocate against the use of this drug during hypothermia. This assumptions gain support by data from in vitro experiments. In isolated rat hearts, both SV and CO were depressed by adrenaline at 28 °C [41], while at 12 °C positive inotropic effects were absent [42]. Interestingly, Schiffmann et al. demonstrated that in normothermic rat hearts, the presence of additionally added calcium would potentiate the inotropic effects of adrenaline. In the hypothermic heart however, increased calcium concentrations [41] mediated a depressive effect of adrenaline on SV and CO. Thus, the negative effects of adrenaline at low temperatures could be a consequence of hypothermia-induced calcium overload, reported in vitro [43], as well as in vivo [29, 30].
### Table 2 β₁-receptor agonists

|                | Adrenaline | Noradrenaline | Isoprenaline | Dobutamine | Dopamine               |
|----------------|------------|---------------|--------------|------------|------------------------|
| **Species**    | Rat, dog   | Cat, dog,     | Rat, dog, rabbit, guinea pig | Dog, pig, rabbit, guinea pig | Dog, pig               |
| **Dosage (in vivo)** | 0.4 μg/kg/min – 4.2 μg/kg/min | 0.2 μg/kg/min – 5.0 μg/kg/min | 5.7 ng/kg/min – 1 μg/kg (bolus) | 2.0 μg/kg/min – 30 μg/kg/min | 2.0 μg/kg/min – 30 μg/kg/min |
| **Administration** | i.v. (in vivo studies), Retrograd coronary perfusion (in vitro studies) | i.v. (in vivo studies) | i.v. (in vivo studies), Retrograd coronary perfusion or in preparation solution (in vitro studies) | i.v. (in vivo studies), Retrograd coronary perfusion or in preparation solution (in vitro studies) | i.v. (in vivo studies) |
| **Target temperature** | 12 °C–28 °C | 28 °C–30 °C | 20 °C–28 °C | 22 °C–31 °C | 25 °C–30 °C |
| **Cardiac effect (hypothermia)** | Elevated CO (low dose) [27, 34, 38]. Depressed CO (high dose) [21, 27, 34, 35, 37]. Negative inotropic effects (in vitro) [41, 42]. | Increased contractile force [26, 45]. Depressed CO [40]. | No effect on CO [50, 51]. Negative or depressed inotropic effect (in vitro) [52, 53]. | Elevated CO [38, 57]. Increased contraction velocity (in vitro) [24]. Reduced or depressed inotropic effect (in vitro) [56]. | Elevated CO or positive inotropic effects [57, 61, 63]. No effect on CO [62, 63]. |
**Noradrenaline**

Noradrenaline has high affinity for $\alpha$-receptors in concert with $\beta_1$-receptor affinity. Infusion of noradrenaline therefore leads both to vasoconstriction of arterioles and a positive inotropic effect [44].

**Noradrenaline during hypothermia** Cotten et al. demonstrated positive inotropic effect of noradrenaline in both normothermic controls and hypothermic (30 °C) dogs [26]. They further described that although this effect was positive, the inotropic effect of noradrenaline was reduced by hypothermia [45]. In cats subjected to moderate hypothermia and rewarming, noradrenaline had a negative effect on CO during hypothermia. The ability of noradrenaline to induce vasoconstriction did however appear intact during cooling, with consistent dose-related increase in MAP [40]. Intact $\alpha$-receptor function is also apparent during noradrenaline infusion in humans cooled on cardiopulmonary bypass to 28–32 °C. In these patients MAP increased significantly with noradrenaline [46]. This can be explained by intact $\alpha_1$-receptor function during a wide range of temperatures, as demonstrated in sheep arteries where vascular response to noradrenaline abolished first when cooling to 5 °C [47]. The sensitivity of $\alpha$-receptors was even found to be increased in human skin artery preparations cooled to 24 °C [48].

**Isoprenaline**

Isoprenaline is a non-selective $\beta$-receptor agonist, lacking $\alpha$-agonist effect. On this background studies looking exclusively at $\beta$-receptor stimulation often use isoprenaline as a model drug. During normothermic conditions, isoprenaline will increase CO and decrease SVR through $\beta_1$- and $\beta_2$-receptor stimulation [49].

**Isoprenaline during hypothermia** Conflicting results exist of pharmacologic effects of isoprenaline at low temperatures, especially when comparing results obtained from in vitro interventional studies of cardiac tissue from different species. Lauri et al. studied in vivo hemodynamic effects of isoprenaline before, during and after severe hypothermia (25 °C) in dogs. Isoprenaline had no positive inotropic effect during hypothermia, but a significant decrease in SVR indicated at least partly intact $\beta_2$-receptor response to stimulation [50], also seen in man at 28–32 °C [46]. Inotropic effects of increasing isoprenaline doses have been investigated in our intact rat model [51] at normothermia and during cooling to 24 °C. We reported that the dose-related positive inotropic response (increase in SV and CO) to isoprenaline at 37 °C were lost during cooling to 24 °C except for the highest dose (20 ng/min). This alteration in response to $\beta$-receptor stimulation also remained after rewarming as only the highest dose of isoprenaline managed to elevate SV above baseline [51]. This finding finds support in another study reporting reduced inotropic response to both isoprenaline and adrenaline in hypothermic rabbit atria at 23 °C [53]. In contrast, in isolated guinea pig hearts cooled to 27 °C isoprenaline still increased the contractility parameter LV dp/dt max, this was accompanied by a similar increase in heart rate, also mediated by $\beta_1$-receptor stimulation [54]. Further, an experiment on isolated atria from guinea pig showed increased inotropic effects of isoprenaline at 25 °C [22]. Sustained ability of adrenaline and isoprenaline to increase contraction amplitude and rate has also been reported when cooling rabbit hearts to 22 °C [24].

**Dobutamine**

Dobutamine predominantly binds to $\beta_1$-receptors and has a weak effect on $\beta_2$- and $\alpha$-receptors. Thus, administering dobutamine in normothermic conditions elevates CO [55].

**Dobutamine during hypothermia** The inotropic effects of dobutamine shows temperature dependency when tested in in vitro guinea pig trabecula. Rieg et al. [56] therefore concluded that elevating cAMP through $\beta_1$-receptor stimulation for providing inotropic support in hypothermic hearts is not an optimal strategy. In isolated rabbit hearts however, dobutamine infusion increased contraction velocity at 22 °C [24]. Increased cardiac output was also observed in response to
dobutamine infusion during hypothermia (30 °C) in pigs [57]. Administering dobutamine in an intact dog model during reperfusion following 60 min global ischemia at 28 °C also showed promising cardiac effects, with increased SV during rewarming using cardio-pulmonary bypass [38].

Dopamine
Like adrenaline, noradrenaline, isoprenaline and dobutamine, dopamine is a catecholamine, giving dose-dependent stimulation of α- and β-receptors and giving a dose-dependent positive inotropic effect in normothermia [58]. Different from selective adrenergic agonists, dopamine also exerts its inotropic and vasoactive effects through stimulation of dopamine receptors [59].

Dopamine during hypothermia The use of dopamine as a vasopressor is recommended in the Up To Date accidental hypothermia guidelines [60]. This is supported by a better cardiovascular recovery in dopamine-treated dogs, after core cooling to 25 °C and subsequent rewarming [61]. Likewise, positive inotropic effects of dopamine were found in pigs core cooled to 30 °C [57]. However, in pigs surface cooled to 32 °C [62] and 25 °C [63] dopamine did not elevate CO. In the latter study, which was conducted in our lab [63], dopamine infusion at 25 °C gave a four-fold increase in plasma concentration compared to normothermia. In difference from other β-adrenergic drugs, cardiovascular responses of dopamine were restored during rewarming to 30 °C [63]. It is therefore apparent from different animal models that dopamine supports cardiac function during rewarming, but it is uncertain whether these positive effects are present during hypothermia below 30 °C. Based on these findings, dopamine is the preferred drug for giving cardiac support during rewarming in the Northern-Norwegian guidelines for accidental hypothermia [64].

Conclusion
Pharmacodynamics
A lack of human studies evaluating cardiovascular effects of adrenergic drugs during hypothermia exists [65]. In the meantime such information can be collected from preclinical experimental studies. This information provide important insight on the effects of pharmacologic interventions already applied in the hypothermic patient. It is apparent that inotropic response to β-adrenergic stimulation seems to be depressed during severe hypothermia. This response is also depressed after rewarming in animals that have received such drugs during hypothermia [51]. In hypothermic animals, adrenaline increases SVR [21]. Studies on dobutamine administration indicates a positive effect, but these studies are carried out at temperatures around 30 °C [38, 57].

Advice of the current guidelines, not to use adrenergic drugs like adrenaline below 30 °C is supported by these preclinical observations, but recommendations for inotropic support in severe accidental hypothermic (>30 °C) patients are needed. Dopamine does not seem to have the same detrimental effects on cardiovascular function in severe accidental hypothermia as adrenaline, even in high plasma concentrations [63], but lack effect before patients are rewarmed to higher temperatures.

Pharmacokinetics
Knowledge about pharmacokinetic effects of adrenergic drugs during hypothermia is limited. However, all pharmacokinetic processes are temperature-dependent; including absorption, distribution and elimination (metabolism and excretion). In general, lowered temperatures slow all these processes down. The time to reach distribution equilibrium will be lengthened and metabolism in the liver and active excretion in the kidneys reduced. Thus half-life (T1/2) of active substances is increased at low temperatures [66]. In humans, it is known that the cytochrome P450 enzyme system is affected by hypothermia. Tortorici [67] found that this resulted in a 7–22% reduced clearance per degree below 37 °C of opiates, barbiturates, benzodiazepines and neuromuscular blockers. Hypothermia does also induce changes in pharmacokinetics of adrenergic drugs. Reduced catechol-O-methyl transferase activity has therefore been suggested to explain a hypothermia-induced hypersensitivity to β-adrenoceptor agonists [24]. Increased T1/2 of adrenaline might therefore have contributed to elevated stimulation of β-adrenoceptors in a recent study reporting increased cAMP levels during 5 min adrenaline administration in hypothermia [21], as the normothermic T1/2 of adrenaline is 2 min [68]. Reduced enzymatic breakdown of cAMP through reduced phosphodiesterase 3 activity, or reduced extracellular release of cAMP as observed in cold fibroblasts [69], might also have boosted adrenaline-mediated cAMP increase in hypothermic hearts [21]. Hypothemia-induced increase in T1/2 is apparent for other catecholamines. At 25 °C in anesthetised pigs, we found that dopamine infusion yielded plasma concentrations 4 times higher than during normothermia. The half-life of dopamine was doubled at this temperature and returned to normothermic values first at 35 °C during rewarming [63]. The high concentrations of dopamine were however not associated with any negative hemodynamic effects.

Apart from temperature-dependent pharmacologic changes in ligand-receptor kinetics, changes in temperature also exert significant alterations in other determinants of cardiac function, which may limit the expected pharmacologic effects achieved at normothermia. During cooling, studies on isolated papillary muscle
show a positive inotropic effect of hypothermia per se [70]. In the intact pig however, cooling induce a reduc-
tion of cardiac contractile function and SV [33]. Lewis
and colleagues showed that the inotropic effect of in-
creasing heart rate during normothermic conditions in
man, is lost at a core temperature of 33 °C [71], inde-
pendent of pharmacologic interventions. Consequently,
hypothermia-induced changes in physiology, not related
to ligand-receptor kinetics, could also be involved in al-
tered pharmacodynamics of β-adrenoceptor agonists
during hypothermic conditions. Important determinants
of blood flow, like blood viscosity, are affected already at
moderate hypothermia [72]. Thus, lack of ability of the
cold blood to increase flow may be part of the chal-
 lenging task to provide positive inotropic, pharmaco-
lologic support during hypothermia. The apparent depressed
function of β1-receptor agonists to provide inotropic ef-
fect in vivo below 30 °C, might therefore be multi-
factorial. Updated guidelines on treatment of hypothermic
patients depend on further studies exploring physio-
logical effects of hypothermia and rewarming, as well as
broader knowledge on the hypothermia-induced changes
on pharmacodynamic and pharmacokinetic effects of
drugs applied in clinical practice.

Clinical implications
Findings in the reviewed literature indicate that negative
or lacking effects of adrenergic drugs during hypothermia
appears to be of multifactorial origin. Our findings from a
majority of pre-clinical studies therefore advocate that
drugs providing adrenergic receptor agonism should be
used carefully during hypothermia and rewarming. Such
information is highly relevant in the clinical setting. Re-
ports show that pharmacological interventions are being
used to provide cardiovascular support in a large propor-
tion of patients during rewarming from accidental
hypothermia [7, 8]. Hypothermia is also used as a ther-
apeutic measure. Comatose survivors of cardiac arrest are
often cooled to temperatures between 32–36 °C for cer-
bral protection [73]. More than 50% of this patient group
are in need of inotropic support to facilitate adequate cir-
culation [74]. Cooling and rewarming of patients down to,
and occasionally below 20 °C is also used for cerebral pro-
tection during procedures like aortic arch surgery [73].
Providing optimal pharmacological, cardiovascular sup-
port in hypothermic patients therefore seems essential,
both in therapeutic hypothermia, and when aiming to im-
prove a high mortality rate associated with accidental
hypothermia [2]. In pigs, dopamine appear a safe way to
provide inotropic support, but lack effect at lower temper-
atures. Further, experimental studies have explored effects
of inotropic pathways like PDE3 inhibition and calcium
sensitizing, drugs that avoid the G-protein coupled
adrenergic receptors. These experiments show promising
results [75–79] on cardiovascular function, both during
cooling and rewarming. However, information from such
pre-clinical studies should be interpreted with care, when
translated to a clinical, human setting. Aiming to provide
better treatment, we call for further studies on physiology,
therapeutic interventions and careful evaluation of ino-
tropic drugs, used in hypothermic patients.

Abbreviations
cAMP: Cyclic AMP; CO: Cardiac output; MAP: Mean arterial pressure;
PKA: Protein kinase A; SV: Stroke volume; SVR: Systemic vascular resistance;
T1/2: Half-life

Acknowledgements
Not applicable.

Funding
The authors’ salaries were funded by their respective affiliations.

Availability of data and materials
Not applicable. All data and material are available in the referenced articles.

Authors’ contributions
Wrote, or contributed to write the manuscript: ESD, GS and TT. All authors
have read and approved the final version of the manuscript.

Competing interests
The authors declare that they have no competing interests.

Consent for publication
Not applicable.

Ethics approval and consent to participate
Not applicable.

Author details
1Anesthesia and Critical Care Research Group, Department of Clinical
Medicine, UiT, The Arctic University of Norway, 9037 Tromsø, Norway.
2Department of Research and Education, Norwegian Air Ambulance
Foundation, 1441 Drabak, Norway. 3Experimental and Clinical Pharmacology,
Department of medical biology, UiT, The Arctic University of Norway, 9037
Tromsø, Norway. 4Division of Surgical Medicine and Intensive Care, University
Hospital of North Norway, 9038 Tromsø, Norway.

Received: 6 October 2016 Accepted: 29 November 2016
Published online: 05 December 2016

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