Inotropes and Vasopressors

Review of Physiology and Clinical Use in Cardiovascular Disease

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Inotropic and vasopressor agents have increasingly become a therapeutic cornerstone for the management of several important cardiovascular syndromes. In broad terms, these substances have excitatory and inhibitory actions on the heart and vascular smooth muscle, as well as important metabolic, central nervous system, and presynaptic autonomic nervous system effects. They are generally administered with the assumption that short- to medium-term clinical recovery will be facilitated by enhancement of cardiac output (CO) or vascular tone that has been severely compromised by often life-threatening clinical conditions. The clinical efficacy of these agents has been investigated largely through examination of their impact on hemodynamic end points, and clinical practice has been driven in part by expert opinion, extrapolation from animal studies, and physician preference. Our aim is to review the mechanisms of action of common inotropes and vasopressors and to examine the contemporary evidence for their use in important cardiac conditions.

Cardiovascular Effects of Common Inotropes and Vasopressors

Catecholamines

Since the initial discovery of epinephrine, the principal active substance from the adrenal gland, the pharmacology and physiology of a large group of endogenous and synthetic catecholamines or “sympathomimetics” have been characterized. Catecholamines mediate their cardiovascular actions predominantly through α, β, and dopaminergic receptors, the density and proportion of which modulate the physiological responses of inotropes and pressors in individual tissues. β-Adrenergic receptor stimulation results in enhanced myocardial contractility through Ca\(^{2+}\)-mediated facilitation of the actin-myosin complex binding with tropo- nin C and enhanced chronicity through Ca\(^{2+}\) channel activation (Figure 1). β-Adrenergic receptor stimulation on vascular smooth muscle cells through a different intracellular mechanism results in increased Ca\(^{2+}\) uptake by the sarcoplasmic reticulum and vasodilation (Figure 1). Activation of α-adrenergic receptors on arterial vascular smooth muscle cells results in smooth muscle contraction and an increase in systemic vascular resistance (SVR; Figure 2). Finally, stimulation of D and D dopaminergic receptors in the kidney and splanchnic vasculature results in renal and mesenteric vasodilation through activation of complex second-messenger systems.

A continuum exists between the effects of the predominantly α-stimulation of phenylephrine (intense vasoconstriction) to the β-stimulation of isoproterenol (marked increase in contractility and heart rate; Table). Specific cardiovascular responses are further modified by reflexive autonomic changes after acute blood pressure alterations, which impact heart rate, SVR, and other hemodynamic parameters. Adrenergic receptors can be desensitized and downregulated in certain conditions, such as in chronic heart failure (HF). Finally, the relative binding affinities of individual inotropes and vasopressors to adrenergic receptors can be altered by hypoxia or acidosis, which mutes their clinical effect.

Dopamine

Dopamine, an endogenous central neurotransmitter, is the immediate precursor to norepinephrine in the catecholamine synthetic pathway (Figure 3A). When administered therapeutically, it acts on dopaminergic and adrenergic receptors to elicit a multitude of clinical effects (Table). At low doses (0.5 to 3 μg·kg\(^{-1}\)·min\(^{-1}\)), stimulation of dopaminergic D receptors postsynaptic receptors concentrated in the coronary, renal, mesenteric, and cerebral beds and D presynaptic receptors present in the vasculature and renal tissues promotes vasodilation and increased blood flow to these tissues. Dopamine also has direct natriuretic effects through its action on renal tubules. The clinical significance of “renal-dose” dopamine is somewhat controversial, however, because it does not increase glomerular filtration rate, and a renal protective effect has not been demonstrated. At intermediate doses (3 to 10 μg·kg\(^{-1}\)·min\(^{-1}\)), dopamine weakly binds to β-adrenergic receptors, promoting norepinephrine release and inhibiting reuptake in presynaptic sympathetic nerve terminals, which results in increased cardiac contractility and chronotropy, with a mild increase in SVR. At higher infusion rates (10 to 20 μg·kg\(^{-1}\)·min\(^{-1}\)), α-adrenergic receptor-mediated vasoconstriction dominates.

Dobutamine

Dobutamine is a synthetic catecholamine with a strong affinity for both β- and β-receptors, which binds to a 3:1 ratio (Table; Figure 3B). With its cardiac β-stimulatory
myocytes, which results in increased concentrations of cAMP. In cardiac stimulatory Gs-GTP unit, activates the adenyl cyclase system, has minimal chronotropic effects, which makes it attractive and has a minimal net impact on CO. Furthermore, this agent primarily increases systolic, diastolic, and pulse pressure with less potent direct inotropic properties. Norepinephrine, the major endogenous neurotransmitter liberated by postganglionic adrenergic nerves (Table; Figure 3A), is a potent α1-adrenergic receptor agonist with modest β-agonist activity, which renders it a powerful vasoconstrictor with less potent direct inotropic properties. Norepinephrine primarily increases systolic, diastolic, and pulse pressure and has a minimal net impact on CO. Furthermore, this agent has minimal chronotropic effects, which makes it attractive.

Epinephrine
Epinephrine is an endogenous catecholamine with high affinity for β1-, β2-, and α1-receptors present in cardiac and vascular smooth muscle (Figure 3A; Table). β-Adrenergic effects are more pronounced at low doses and α1-adrenergic effects at higher doses. Coronary blood flow is enhanced through an increased relative duration of diastole at higher heart rates and through stimulation of myocytes to release local vasodilators, which largely counterbalance direct α1-mediated vasoconstriction. Arterial and venous pulmonary pressures are increased through direct pulmonary vasoconstriction and increased pulmonary blood flow. High and prolonged doses can cause direct cardiac toxicity through damage to arterial walls, which causes focal regions of myocardial contraction band necrosis, and through direct stimulation of myocyte apoptosis.

Isoproterenol
Isoproterenol is a potent, nonselective, synthetic β-adrenergic agonist with very low affinity for α-adrenergic receptors for use in settings in which heart rate stimulation may be undesirable. Coronary flow is increased owing to elevated diastolic blood pressure and indirect stimulation of cardiac myocytes, which release local vasodilators. Prolonged nor- epinephrine infusion can have a direct toxic effect on cardiac myocytes by inducing apoptosis via protein kinase A activation and increased cytosolic Ca2+ influx.

Figure 1. Simplified schematic of postulated intracellular actions of β-adrenergic agonists. β-Receptor stimulation, through a stimulatory Gs-GTP unit, activates the adenyl cyclase system, which results in increased concentrations of cAMP. In cardiac myocytes, β-receptor activation through increased cAMP concentration activates Ca2+ channels, which leads to Ca2+-mediated enhanced chronotropic responses and positive inotropy by increasing the contractility of the actin-myosin-troponin system. In vascular smooth muscle, β-stimulation and increased cAMP results in stimulation of a cAMP-dependent protein kinase, phosphorylation of phospholamban, and augmented Ca2+ uptake by the sarcoplasmic reticulum (SR), which leads to vasodilation. Adapted from Gillies et al with permission of the publisher.

Figure 2. Schematic representation of postulated mechanisms of intracellular action of α1-adrenergic agonists. α1-Receptor stimulation activates a different regulatory G protein (Gq), which acts through the phospholipase C system and the production of 1,2-diacylglycerol (DAG) and, via phosphatidylinositol-4,5-biphosphate (PiP2), of inositol 1,4,5-triphosphate (IP3). IP3 activates the release of Ca2+ from the sarcoplasmic reticulum (SR), which by itself and through Ca2+-calmodulin-dependent protein kinases influences cellular processes, leading in vascular smooth muscle to vasoconstriction. Adapted from Gillies et al with permission of the publisher.
### Table. Inotropic and Vasopressor Drug Names, Clinical Indication for Therapeutic Use, Standard Dose Range, Receptor Binding (Catecholamines), and Major Clinical Side Effects

<table>
<thead>
<tr>
<th>Drug</th>
<th>Clinical Indication</th>
<th>Dose Range</th>
<th>Receptor Binding</th>
<th>Major Side Effects</th>
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<tbody>
<tr>
<td><strong>Catecholamines</strong></td>
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<tr>
<td>Dopamine</td>
<td>Shock (cardiogenic, vasodilatory) HF</td>
<td>2.0 to 20 µg · kg⁻¹ · min⁻¹ (max 50 µg · kg⁻¹ · min⁻¹)</td>
<td>++ + +</td>
<td>Severe hypertension (especially in patients taking nonselective β-blockers)</td>
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<tr>
<td></td>
<td>Symptomatic bradycardia unresponsive to atropine or pacing</td>
<td></td>
<td></td>
<td>Ventricular arrhythmias</td>
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<td></td>
<td></td>
<td></td>
<td></td>
<td>Cardiac ischemia</td>
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<td></td>
<td></td>
<td>Tissue ischemia/gangrene (high doses or due to tissue extravasation)</td>
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<tr>
<td>Dobutamine</td>
<td>Low CO (decompensated HF, cardiogenic shock, sepsis-induced myocardial dysfunction)</td>
<td>2.0 to 20 µg · kg⁻¹ · min⁻¹ (max 40 µg · kg⁻¹ · min⁻¹)</td>
<td>++ + + +</td>
<td>Tachycardia</td>
</tr>
<tr>
<td></td>
<td>Symptomatic bradycardia unresponsive to atropine or pacing</td>
<td></td>
<td></td>
<td>Increased ventricular response rate in patients with atrial fibrillation</td>
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<td></td>
<td>Ventricular arrhythmias</td>
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<td></td>
<td>Cardiac ischemia</td>
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<td></td>
<td></td>
<td>Hypertension (especially nonselective β-blocker patients)</td>
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<td>Hypotension</td>
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<tr>
<td>Norepinephrine</td>
<td>Shock (vasodilatory, cardiogenic)</td>
<td>0.01 to 3 µg · kg⁻¹ · min⁻¹</td>
<td>+ + + +</td>
<td>Arrhythmias</td>
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<td>Bradycardia</td>
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<td>Peripheral (digital) ischemia</td>
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<tr>
<td>Epinephrine</td>
<td>Shock (cardiogenic, vasodilatory) Cardiac arrest</td>
<td>Infusion: 0.01 to 0.10 µg · kg⁻¹ · min⁻¹</td>
<td>++ + + +</td>
<td>Ventricular arrhythmias</td>
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<td></td>
<td>Bronchospasm/anaphylaxis</td>
<td>Bolus: 1 mg IV every 3 to 5 min (max 0.2 mg/kg)</td>
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<td>Severe hypertension resulting in cerebrovascular hemorrhage</td>
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<td></td>
<td>Symptomatic bradycardia or heart block unresponsive to atropine or pacing</td>
<td>IM: (1:1000): 0.1 to 0.5 mg (max 1 mg)</td>
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<td>Cardiac ischemia</td>
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<td></td>
<td></td>
<td>Sudden cardiac death</td>
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<tr>
<td>Isoproterenol</td>
<td>Bradyarrhythmias (especially torsade des pointes) Brugada syndrome</td>
<td>2 to 10 µg/min</td>
<td>0 + + + + +</td>
<td>Ventricular arrhythmias</td>
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<td></td>
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<td></td>
<td></td>
<td>Cardiac ischemia</td>
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<td></td>
<td></td>
<td>Hypertension</td>
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<td></td>
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<td></td>
<td></td>
<td>Tissue necrosis with extravasation</td>
</tr>
<tr>
<td>Phenylephrine</td>
<td>Hypotension (vagally mediated, medication-induced) Increase MAP with AS and hypotension Decrease LVOT gradient in HCM</td>
<td>Bolus: 0.1 to 0.5 mg IV every 10 to 15 min</td>
<td>+ + + + + 0</td>
<td>Reflex bradycardia</td>
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<td></td>
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<td>Infusion: 0.4 to 9.1 µg · kg⁻¹ · min⁻¹</td>
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<td>Hypertension (especially with nonselective β-blockers)</td>
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<td></td>
<td>Severe peripheral and visceral vasoconstriction</td>
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<td>PDIs</td>
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<td>Tissue necrosis with extravasation</td>
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<tr>
<td>Milrinone</td>
<td>Low CO (decompensated HF, after cardiotomy)</td>
<td>Bolus: 50 µg/kg bolus over 10 to 30 min</td>
<td>N/A</td>
<td>Ventricular arrhythmias</td>
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<td></td>
<td></td>
<td>Infusion: 0.375 to 0.75 µg · kg⁻¹ · min⁻¹ (dose adjustment necessary for renal impairment)</td>
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<td>Hypotension</td>
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<td>Cardiac ischemia</td>
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<td>Torsade des pointes</td>
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<tr>
<td>Amrinone</td>
<td>Low CO (refractory HF)</td>
<td>Bolus: 0.75 mg/kg over 2 to 3 min</td>
<td>N/A</td>
<td>Arrhythmias, enhanced AV conduction</td>
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<td></td>
<td></td>
<td>Infusion: 5 to 10 µg · kg⁻¹ · min⁻¹</td>
<td></td>
<td>(increased ventricular response rate in atrial fibrillation)</td>
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<td></td>
<td></td>
<td></td>
<td></td>
<td>Hypotension</td>
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<td></td>
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<td></td>
<td></td>
<td>Thrombocytopenia</td>
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<td>Hepatotoxicity</td>
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<tr>
<td>Vasopressin</td>
<td>Shock (vasodilatory, cardiogenic) Cardiac arrest</td>
<td>Infusion: 0.01 to 0.1 U/min (common fixed dose 0.04 U/min)</td>
<td>V₁ receptors (vascular smooth muscle)</td>
<td>Arhythmias</td>
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<tr>
<td></td>
<td></td>
<td>Bolus: 40-IV bolus</td>
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<td>Hypertension</td>
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<td></td>
<td>Decreased CO (at doses &gt;0.4 U/min)</td>
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<td>Cardiac ischemia</td>
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<td>Severe peripheral vasoconstriction causing ischemia (especially skin)</td>
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<td>Splanchic vasoconstriction</td>
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<tr>
<td>Levosimendan</td>
<td>Decompensated HF</td>
<td>Loading dose: 12 to 24 µg/kg over 10 min</td>
<td>N/A</td>
<td>Tachycardia, enhanced AV conduction</td>
</tr>
<tr>
<td></td>
<td></td>
<td>Infusion: 0.05 to 0.2 µg · kg⁻¹ · min⁻¹</td>
<td></td>
<td>Hypotension</td>
</tr>
</tbody>
</table>

α₁ indicates α-1 receptor; β₁, β-1 receptor; β₂, β-2 receptor; DA, dopamine receptors; 0, zero significant receptor affinity; + through +++++, minimal to maximal relative receptor affinity; N/A, not applicable; IV, intravenous; IM, intramuscular; max, maximum; AS, aortic stenosis; LVOT, LV outflow tract; HCM, hypertrophic cardiomyopathy; and AV, atriointricular.
Phenylephrine

With its potent synthetic α-adrenergic activity and virtually no affinity for β-adrenergic receptors (Table; Figure 3B), phenylephrine is used primarily as a rapid bolus for immediate correction of sudden severe hypotension. It can be used to raise mean arterial pressure (MAP) in patients with severe hypotension and concomitant aortic stenosis, to correct hypotension caused by the simultaneous ingestion of sildenafil and nitrates, to decrease the outflow tract gradient in patients with obstructive hypertrophic cardiomyopathy, and to correct vagally mediated hypotension during percutaneous diagnostic or therapeutic procedures. This agent has virtually no direct heart rate effects, although it has the potential to induce significant baroreceptor-mediated reflex rate responses after rapid alterations in MAP.

Phosphodiesterase Inhibitors

Phosphodiesterase 3 is an intracellular enzyme associated with the sarcoplasmic reticulum in cardiac myocytes and vascular smooth muscle that breaks down cAMP into AMP. Phosphodiesterase inhibitors (PDIs) increase the level of cAMP by inhibiting its breakdown within the cell, which leads to increased myocardial contractility (Figure 4). These agents are potent inotropes and vasodilators and also improve
diastolic relaxation (lusitropy), thus reducing preload, afterload, and SVR.

Milrinone is the PDI most commonly used for cardiovascular indications (Table). In its parenteral form, it has a longer half-life (2 to 4 hours) than many other inotropic medications. This drug is particularly useful if adrenergic receptors are downregulated or desensitized in the setting of chronic HF, or after chronic β-agonist administration. Amrinone is used less often because of important side effects, which include dose-related thrombocytopenia.

**Vasopressin**
Isolated in 1951, the nonapeptide vasopressin or “antidiuretic hormone” is stored primarily in granules in the posterior pituitary gland and is released after increased plasma osmolality or hypotension, as well as pain, nausea, and hypoxia. Vasopressin is synthesized to a lesser degree by the heart in response to elevated cardiac wall stress and by the adrenal gland in response to increased catecholamine secretion. It exerts its circulatory effects through V1 (V1a in vascular smooth muscle, V1b in the pituitary gland) and V2 receptors (renal collecting duct system; Table). V1a stimulation mediates constriction of vascular smooth muscle, whereas V2 receptors mediate water reabsorption by enhancing renal collecting duct permeability.

Vasopressin causes less direct coronary and cerebral vasomotor constriction than catecholamines and has a neutral or inhibitory impact on CO, depending on its dose-dependent increase in SVR and the reflexive increase in vagal tone. A vasopressin-modulated increase in vascular sensitivity to norepinephrine further augments its pressor effects. The agent may also directly influence mechanisms involved in the pathogenesis of vasodilation, through inhibition of ATP-activated potassium channels, attenuation of nitric oxide production, and reversal of adrenergic receptor downregulation. Thepressor effects of vasopressin are relatively preserved during hypoxic and acidic conditions, which commonly develop during shock of any origin.

**Calcium-Sensitizing Agents**
Calcium sensitizers are a recently developed class of inotropic agents, levosimendan being the most well known (Table). These agents have a dual mechanism of action that includes calcium sensitization of contractile proteins and the opening of ATP-dependent potassium (K⁺) channels. Calcium-dependent binding to troponin C enhances ventricular contractility without increasing intracellular Ca²⁺ concentration or compromising diastolic relaxation, which may favorably impact myocardial energetics relative to traditional inotropic therapies. The opening of K⁺ channels on vascular smooth muscle leads to arteriolar and venous vasodilation and may confer a degree of myocardial protection during ischemia. The combination of improved contractile performance and vasodilation is particularly beneficial during acute and chronic HF states, for which levosimendan is being used with increasing frequency in some countries.

**Evidence for Use of Inotropes and Vasopressors in Cardiovascular Disease**

**Cardiogenic Shock Complicating Acute Myocardial Infarction**
Inotropes and vasopressors are used routinely in the setting of cardiogenic shock complicating acute myocardial infarction (AMI). These agents all increase myocardial oxygen consumption and can cause ventricular arrhythmias, contraction-band necrosis, and infarct expansion. However, critical hypotension itself compromises myocardial perfusion, leading to elevated left ventricular (LV) filling pressures, increased myocardial oxygen requirements, and further reduction in the coronary perfusion gradient. Thus, hemodynamic benefits usually outweigh specific risks of inotropic therapy when used as a bridge to more definitive treatment measures.

Inotropic agents may improve mitochondrial function in noninfarcted myocardium that has become deranged during AMI complicated by shock. However, free cytosolic Ca²⁺, which is significantly elevated in postischemic cardiac myocytes, is further increased with the administration of dopamine, which leads to activation of proteolytic enzymes, proapoptotic signal cascades, mitochondrial damage, and eventual membrane disruption and necrosis. Thus, the lowest possible doses of inotropic and pressor agents should be used to adequately support vital tissue perfusion while limiting adverse consequences, some of which may not be immediately apparent.

The American College of Cardiology/American Heart Association guidelines for management of hypotension complicating AMI suggest the use of dobutamine as a first-line agent if systolic blood pressure ranges between 70 and 100 mm Hg in the absence of signs and symptoms of shock. Dopamine is suggested in patients who have the same systolic blood pressure in the presence of symptoms of shock. However, definitive evidence supporting use of specific agents in this setting is lacking. Moderate doses of these agents maximize inotropy and avoid excessive α₁-adrenergic stimulation that can result in end-organ ischemia. The deliberate combination of dopamine and dobutamine at a dose of 7.5 μg·kg⁻¹·min⁻¹ each was shown to improve hemodynamics and limit important side effects compared with either individual agent administered at 15 μg·kg⁻¹·min⁻¹. Moderate doses of combinations of medications may potentially be more effective than maximal doses of any individual agent.

When response to a medium dose of dopamine or dobutamine in combination is inadequate, or the patient’s presenting systolic blood pressure is <70 mm Hg, the use of norepinephrine has been recommended. With an antithrombotic effect in addition to its pressor qualities, norepinephrine may be the optimal choice under these conditions compared with epinephrine, which can exacerbate lactic acidosis and promote thrombosis in coronary vasculature.

During early shock, endogenous vasopressin levels are elevated significantly to help maintain end-organ perfusion. As the shock state progresses, however, plasma vasopressin levels fall dramatically, which contributes to a loss of...
vascular tone, worsening hypotension, and end-organ perfusion. Proposed mechanisms to explain this phenomenon include depletion of neurohypophyseal stores, baroreceptor and generalized autonomic dysfunction during prolonged shock, and endogenous norepinephrine-induced inhibition of vasopressin release. Vasopressin therapy may thus be effective in norepinephrine-resistant vasodilatory shock, improving MAP, cardiac index, and LV stroke work index and reducing the need for norepinephrine, resulting in decreased cardiotoxicity and malignant arrhythmias. Vasopressin may also attenuate interleukin-induced generation of nitric oxide, have a modest inotropic effect on the myocardium via $V_1_r$-mediated increases in intracellular Ca$^{2+}$, and improve coronary blood flow due to catecholamine sparing.

In the only study to date that examined vasopressin use in cardiogenic shock after AMI, this agent was found to increase MAP without adversely impacting cardiac index and wedge pressure. Cardiac power index, an important determinant of outcome in cardiogenic shock after AMI, was not adversely affected by vasopressin but decreased when norepinephrine was used. Further studies to validate the use of vasopressin in this setting are needed.

**Congestive HF**

Inotropic therapy is used in the management of decompensated HF to lower end-diastolic pressure and improve diuresis, thus allowing traditional medical therapy (eg, angiotensin-converting enzyme inhibitors, diuretics, and $\beta$-blockers) to be re instituted gradually. Patients with decompensated HF unresponsive to diuretics often have diminished concomitant peripheral perfusion, clinically apparent as cool extremities, narrowed pulse pressure, and worsening renal function. They may have markedly elevated SVR despite hypotension due to the stimulation of the renal-angiotensin-aldosterone system, as well as release of endogenous catecholamines and vasopressin. In this setting, reversal of systemic vasoconstriction is often achieved through the use of vasodilators (such as sodium nitroprusside) and inotropes with peripheral vasodilatory properties to improve hemodynamic parameters and clinical symptoms.

The use of positive inotropes (parenteral inotropes and oral PDIs) in chronic HF has been consistently demonstrated to increase mortality. A proposed central mechanism involves a chronic increase in intracellular Ca$^{2+}$, which contributes to altered gene expression and apoptosis and an increased likelihood of malignant ventricular arrhythmias. As a result, the current American College of Cardiology/American Heart Association guidelines for diagnosis and management of chronic HF in the adult do not recommend the routine use of intravenous inotropic agents for patients with refractory end-stage HF (class III recommendation) but do state that they may be considered for palliation of symptoms in these patients (class IIb recommendation). The European Society of Cardiology acute HF guidelines also stress that few controlled trials with intravenous inotropic agents have been performed. However, these guidelines do point out that in an appropriate clinical setting of hypotension and peripheral hyperperfusion, particular agents may be indicated with slightly different levels of recommendation (dobutamine and levosimendan, class IIa; PDIs and dopamine, class IIb).

The most commonly recommended initial inotropic therapies for refractory HF (dobutamine, dopamine, and milrinone) are used to improve CO and enhance diuresis by improving renal blood flow and decreasing SVR without exacerbating systemic hypotension. Dobutamine stimulation of $\beta_1$- and $\beta_2$-receptors can achieve this goal at low to medium doses by modestly increasing contractility with usually mild systemic vasodilation. Unfortunately, $\beta$-adrenergic receptor responses are often blunted in the failing human heart. A chronic increase in activation of the sympathetic nervous system and increased circulating catecholamine levels results in a phosphorylation signal that leads to uncoupling of the surface receptor from its intracellular signal transduction proteins (desensitization), as well as increased receptor targeting for endocytosis (decreased receptor density). PDIs such as milrinone, acting through a non-$\beta$-adrenergic mechanism, are not associated with diminished efficacy or tolerance with prolonged use. This drug causes relatively more significant right ventricular afterload reduction through pulmonary vasodilation and less direct cardiac inotropy, which results in less myocardial oxygen consumption. Milrinone can cause severe systemic hypotension, necessitating the coadministration of additional pressor therapies. Direct randomized comparisons of milrinone and dobutamine have been small and have demonstrated similar clinical outcomes.

Several major clinical trials have evaluated the safety and efficacy of levosimendan in HF syndromes. Two early studies demonstrated a mortality benefit in patients given levosimendan versus placebo early (within 14 days) in the setting of LV failure complicating AMI (RUSSLAN [Randomized Study on Safety and Effectiveness of Levosimendan in Patients With Left Ventricular Failure due to an Acute Myocardial Infarct]) and at 180 days in the setting of chronic HF when compared with dobutamine therapy (LIDO [Levosimendan Infusion versus Dobutamine in Severe Low-Output Heart Failure]). However, in larger multicenter randomized trials in the setting of acute decompensated HF (REVIVE II [Randomized Multicenter Evaluation of Intravenous Levosimendan Efficacy] and SURVIVE [Survival of Patients With Acute Heart Failure in Need of Intravenous Inotropic Support]), le vosimendan use significantly improved symptoms but not survival.

In some patients, complete inotropic dependence manifested by symptomatic hypotension, recurrent congestive symptoms, or worsening renal function may develop after discontinuation of parenteral therapy. Inotropic support may become necessary until cardiac transplantation or implantation of an LV assist device can be instituted. Long-term therapy is also used as a “bridge to decision” in patients who are not presently destination-therapy candidates but may become so in the future. Inotrope-dependent HF patients who do not go on to definitive therapy have a poor prognosis, with 1-year mortality ranging from 79% to 94%. Long-term inotropic therapy is associated with an increased risk of line sepsis, arrhythmias, accelerated functional decline due to worsening nutritional status, and direct acceleration of end-
organ dysfunction, such as the development of cosinophilic myocarditis from an allergic response to chronic dobutamine exposure. Inotropic home therapy has been used effectively for palliation of symptoms in patients who are not candidates for LV assist device support or transplantation, enabling those individuals to die in the comfort of their own homes.

The majority of HF patients can be weaned off inotropic infusions successfully after diuresis of excess volume and careful adjustment of concomitant oral medications. General recommendations have been to keep patients in the hospital and on a stable oral HF regimen for 48 hours before discharge to ensure adequacy of the initiated therapy.

Cardiopulmonary Arrest
Inotropic and vasopressor agents are a mainstay of resuscitation therapy during cardiopulmonary arrest. Epinephrine, with its potent vasopressor and inotropic properties, can rapidly increase diastolic blood pressure to facilitate coronary perfusion and help restore organized myocardial contractility. However, it is not clear whether epinephrine actually facilitates cardioversion to normal rhythm, and its use has been associated with increased oxygen consumption, ventricular arrhythmias, and myocardial dysfunction after successful resuscitation. Repeated high-bolus doses (5 mg) appear no more effective than repeated standard doses (1 mg) at restoring circulation.

The finding that endogenous vasopressin levels are greater in patients successfully resuscitated from sudden cardiac death than in nonsurvivors sparked interest in the use of vasopressin for this indication. Experimentally, the use of vasopressin during cardiopulmonary collapse has demonstrated a more beneficial effect than epinephrine on cerebral and myocardial blood flow, resulting in more sustained increases in MAP. Clinically, its use has been associated with a higher rate of short-term survival in patients experiencing out-of-hospital ventricular fibrillation. However, in a larger trial of 1186 patients with out-of-hospital cardiac arrest who were randomized to 2 injections of either 40 U of vasopressin or 1 mg of epinephrine (followed by additional treatment with epinephrine if needed), patients with asystole but not those with ventricular fibrillation or pulseless electrical activity were significantly more likely to survive to hospital admission with vasopressin administration. The mechanism of benefit may stem from the ability of vasopressin to retain its potent vasoconstricting properties under severely acidic conditions, in which catecholamines have limited efficacy. The current American Heart Association guidelines for adult cardiac life support have incorporated vasopressin as a 1-time alternative to the first or second dose of epinephrine (1-time bolus of 40 U) in patients with pulseless electrical activity or asystole and for pulseless ventricular tachycardia or ventricular fibrillation.

Postoperative Cardiac Surgery
Pharmacological support may be necessary during and after weaning from cardiopulmonary bypass in patients who have developed a low-CO syndrome, arbitrarily defined as a cardiac index <2.4 L·min⁻¹·m⁻² with evidence of end-organ dysfunction. Causes of low CO include cardioplegia-induced myocardial dysfunction, precipitation of cardiac ischemia during aortic cross-clamping, reperfusion injury, activation of inflammatory and coagulation cascades, and the presence of nonrepaired preexisting cardiac disease. Therapy should be instituted promptly in addition to other measures, including optimization of volume status, reduction of SVR with propofol infusion, temporary pacing, and intra-aortic balloon counterpulsation. Although no single agent is universally superior in this setting, dobutamine has the most desirable side-effect profile of the β-agonists, whereas PDIs increase flow through arterial grafts, reduce MAP, and improve right-sided heart performance in pulmonary hypertension. As is the case in HF, concomitant vasopressor therapy may be necessary.

Several studies have examined the role of prophylactic inotropic or vasopressor therapy in weaning from cardiopulmonary bypass or to improve hemodynamic status in general. Preemptive milrinone administration before separation from cardiopulmonary bypass was found to attenuate postoperative deterioration in cardiac function and reduce the need for additional inotropes. In off-pump bypass surgery patients, the use of preemptive milrinone significantly ameliorates increases in mitral regurgitation and improves hemodynamic indexes that often deteriorate with off-pump surgery. Milrinone and dobutamine were both found to be effective in improving general hemodynamic parameters compared with placebo in a European multicenter, randomized, open-label trial. The development of a systemic inflammatory response during cardiopulmonary bypass may cause severe generalized vasodilation, known as “vasoplegia syndrome,” which can result in increased early mortality, especially in heart transplant recipients. This syndrome is associated with prolonged cardiopulmonary bypass time, orthotropic heart transplantation, and LV assist device insertion and is characterized by severe persistent hypotension, metabolic acidosis, decreased SVR, and low intracardiac filling pressures, with normal or elevated CO. Preoperative risk factors include preoperative angiotensin-converting enzyme inhibitor, calcium channel blocker, or intravenous heparin use and poor LV function. Development of vasoplegia syndrome may be related to the release of vasodilatory inflammatory mediators, extensive complement activation, or vasoactive substance depletion, such as vasopressin. Although catecholamine therapy is often ineffective, methylene blue (through a nitric oxide–inhibition mechanism) and vasopressin have been shown to improve outcomes.

Right Ventricular Infarction
Significant right ventricular free-wall ischemia leads to immediate dilation of the right ventricle within a constrained pericardium. A rapid increase in intrapericardial pressure and intraventricular septal shift alters LV geometry, impairing LV filling and contractile performance. These combined effects result in a drop in CO that may exacerbate shock. Excessive intravenous fluid beyond a right atrial pressure >15 mm Hg to improve a “preload-dependent” right ventricle can result in deterioration of LV performance. Dobutamine improves myocardial performance in this setting.
Close observation is essential to monitor for exacerbation of hypotension and atrial arrhythmias, which can profoundly worsen hemodynamics.

**Bradyarrhythmias**

Owing to their chronotropic effects, β-adrenergic agonists can be useful for transient emergency treatment of bradyarrhythmias if atropine is ineffective. The use of the β-agonists dobutamine, dopamine, or isoproterenol can stabilize the patient to allow time for a temporary pacemaker to be inserted. These agents are also useful under the same circumstances to treat bradycardia-induced torsade des points. Finally, isoproterenol has also been used to suppress the trigger for ventricular fibrillation in patients with the Brugada syndrome who do not wish to have cardioverter-defibrillator implantation to prevent sudden cardiac death.

**Adjuvant Issues**

**Patient Monitoring During Parenteral Inotropic and Vasopressor Therapy**

Patients requiring treatment with inotropes and vasopressors generally require monitoring in an intensive care or step-down setting because of the potential for development of life-threatening arrhythmias.

**Invasive Blood Pressure Monitoring**

In shock, continuous blood pressure monitoring with an arterial line is essential both to monitor the status of the underlying illness and because inotropes and vasopressors have the potential to induce life-threatening hypotension or hypertension. Chronic HF patients undergoing hemodynamic tailoring with a low-dose β-agonist or PDI can usually be monitored noninvasively.

**Pulmonary Artery Catheter Use**

Consensus on pulmonary artery catheter use during treatment with inotropic therapy is lacking. Although this tool can be helpful diagnostically, its routine use has never been shown to improve outcomes. This may reflect an absence of effective evidence-based therapies to be used in response to pulmonary artery catheter data in the treatment of critically ill patients.

In the ESCAPE trial (Evaluation Study of Congestive Heart Failure and Pulmonary Artery Catheterization Effectiveness), which examined pulmonary artery catheter use in patients with severe HF, catheter insertion was deemed safe but was not associated with improved rates of mortality or hospitalization. Inotropic titration with pulmonary artery catheter data in isolation can result in inappropriate stimulation of CO, thus negatively impacting prognosis in heterogeneous intensive care unit patient populations. Titration of inotropic therapy should be guided by the adequacy of end-organ perfusion, based on multiple clinical parameters.

**Goals of Inotropic and Vasopressor Therapy**

The use of inotropes and vasopressors has not been shown in randomized, controlled studies to ultimately lead to improved patient outcomes, at least in part because no clinical trials have been conducted with study size and power adequate to test their effect on improving survival. In the absence of such data, the definitive goals of therapy must be considered of primary importance, and the role of inotropic therapy should be kept in a supportive context to allow treatment of the underlying disorder. Such therapy includes prompt percutaneous or surgical revascularization and the institution of mechanical support (intra-aortic balloon counterpulsation or LV assist device) to improve coronary perfusion, CO, or both.

**Conclusions and Recommendations**

In conclusion, inotropes and vasopressors play an essential role in the supportive care of a number of important cardiovascular disease processes. To date, prospective examination of their impact on clinical outcomes in randomized trials has been minimal, despite their widespread use in cardiovascular illness. However, the recently published TRIUMPH (Tilarginine Acetate Injection in a Randomized International Study in Unstable MI Patients With Cardiogenic Shock) international randomized trial of N6-monomethyl L-arginine in cardiogenic shock has shown that such trials are not only feasible but necessary to validate findings of smaller studies. A better understanding of the physiology and important adverse effects of these medications should lead to directed clinical use, with realistic therapeutic goals. The following broad recommendations can be made:

Smaller combined doses of inotropes and vasopressors may be advantageous over a single agent used at higher doses to avoid dose-related adverse effects. The use of vasopressin at low to moderate doses may allow catecholamine sparing, and it may be particularly useful in settings of catecholamine hyposensitivity and after prolonged critical illness. In cardiogenic shock complicating AMI, current guidelines based on expert opinion recommend dopamine or dobutamine as first-line agents with moderate hypotension (systolic blood pressure 70 to 100 mm Hg) and norepinephrine as the preferred therapy for severe hypotension (systolic blood pressure <70 mm Hg).

Routine inotropic use is not recommended for end-stage HF. When such use is essential, every effort should be made to either reinstitute stable oral therapy as quickly as possible or use destination therapy such as cardiac transplantation or LV assist device support. Large randomized trials focusing on clinical outcomes are needed to better assess the clinical efficacy of these agents.

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