478e-2 TABLE 478e-2 PHYSIOLOGIC CHANGES ASSOCIATED WITH ACCIDENTAL HYPOTHERMIA

Covority	Podu Tomporaturo	Central Nervous	Cardiovaccular	Decreivatory	Donal and Endocrino	Nouromuccular
Severity		System	Tachycardia, than pro	Tachypped then	Renal and Endocrine	Increased precisiver
Mild	55 € (95 F)− 52.2 € (90°F)	cerebral metabolism; amnesia; apathy; dysar- thria; impaired judgment; maladaptive behavior	gressive bradycardia; cardiac cycle prolonga- tion; vasoconstriction; increase in cardiac out- put and blood pressure	progressive decrease in respiratory minute volume; declining oxygen consumption; bronchorrhea; bron- chospasm	catecholamines, adre- nal steroids, triiodothy- ronine, and thyroxine; increase in metabolism with shivering	ing muscle tone, then fatiguing
Moderate	<32.2°C (90°F)–28°C (82.4°F)	EEG abnormalities; pro- gressive depression of level of consciousness; pupillary dilation; para- doxical undressing; hal- lucinations	Progressive decrease in pulse and cardiac output; increased atrial and ventricular arrhyth- mias; suggestive (J- wave) ECG changes	Hypoventilation; 50% decrease in carbon dioxide production per 8°C (46°F) drop in temperature; absence of protective airway reflexes	50% increase in renal blood flow; renal autoregulation intact; impaired insulin action	Hyporeflexia; dimin- ishing shivering- induced thermogen- esis; rigidity
Severe	<28°C (<82.4°F)	Loss of cerebrovascular autoregulation; decline in cerebral blood flow; coma; loss of ocular reflexes; progressive decrease in EEG abnor- malities	Progressive decrease in blood pressure, heart rate, and cardiac out- put; reentrant dysrhyth- mias; maximal risk of ventricular fibrillation; asystole	Pulmonic conges- tion and edema; 75% decrease in oxygen consumption; apnea	Decrease in renal blood flow that parallels decrease in cardiac output; extreme oli- guria; poikilothermia; 80% decrease in basal metabolism	No motion; decreased nerve- conduction velocity; peripheral areflexia; no corneal or oculo- cephalic reflexes

Abbreviations: ECG, electrocardiogram; EEG, electroencephalogram.

Source: Modified from DF Danzl, RS Pozos: N Engl J Med 331:1756, 1994.

When a patient in hypothermic cardiac arrest is first discovered, cardiopulmonary resuscitation is indicated unless (1) a do-not-resuscitate status is verified, (2) obviously lethal injuries are identified, or (3) the depression of a frozen chest wall is not possible. As the resuscitation proceeds, the prognosis is grave if there is evidence of widespread cell lysis, as reflected by potassium levels >10–12 mmol/L (10–12meq/L). Other findings that may preclude continuing resuscitation include a core temperature <10–12°C (<50–54°F), a pH <6.5, and evidence of intravascular thrombosis with a fibrinogen value <0.5 g/L (<50 mg/ dL). The decision to terminate resuscitation before rewarming the patient past 33°C (91°F) should be predicated on the type and severity of the precipitants of hypothermia. There are no validated prognostic indicators for recovery from hypothermia. A history of asphyxia with secondary cooling is the most important negative predictor of survival.

DIAGNOSIS AND STABILIZATION

Hypothermia is confirmed by measurement of the core temperature, preferably at two sites. Rectal probes should be placed to a depth of 15 cm and not adjacent to cold feces. A simultaneous esophageal probe should be placed 24 cm below the larynx; it may read falsely high during heated inhalation therapy. Relying solely on infrared tympanic thermography is not advisable.

After a diagnosis of hypothermia is established, cardiac monitoring should be instituted, along with attempts to limit further heat loss. If the patient is in ventricular fibrillation, it is unclear at what core temperature ventricular defibrillation (2 J/kg) should first be attempted. One attempt below 30°C is warranted. Further defibrillation attempts should be deferred until some rewarming $(1^{\circ}-2^{\circ}C)$ is achieved and ventricular fibrillation is coarser. Although cardiac pacing for hypothermic bradydysrrhythmias is rarely indicated, the transthoracic technique is preferable.

Supplemental oxygenation is always warranted, since tissue oxygenation is affected adversely by the leftward shift of the oxyhemoglobin dissociation curve. Pulse oximetry may be unreliable in patients with vasoconstriction. If protective airway reflexes are absent, gentle endotracheal intubation should be performed. Adequate preoxygenation will prevent ventricular arrhythmias.

Insertion of a gastric tube prevents dilation secondary to decreased bowel motility. Indwelling bladder catheters facilitate monitoring of cold-induced diuresis. Dehydration is encountered commonly with chronic hypothermia, and most patients benefit from an intravenous or intraosseous bolus of crystalloid. Normal saline is preferable to lactated Ringer's solution, as the liver in hypothermic patients inefficiently metabolizes lactate. The placement of a pulmonary artery catheter can cause perforation of the less compliant pulmonary artery. Insertion of a central venous catheter deeply into the cold right atrium should be avoided since this procedure can precipitate arrhythmias.

Arterial blood gases should not be corrected for temperature (Chap. 66). An uncorrected pH of 7.42 and a Pco_2 of 40 mmHg reflect appropriate alveolar ventilation and acid-base balance at any core temperature. Acid-base imbalances should be corrected gradually, since the bicarbonate buffering system is inefficient. A common error is overzealous hyperventilation in the setting of depressed CO_2 production. When the Pco_2 decreases by 10 mmHg at 28°C (82°F), it doubles the pH increase of 0.08 that occurs at 37°C (99°F).

The severity of anemia may be underestimated because the hematocrit increases 2% for each 1°C drop in temperature. White blood cell sequestration and bone marrow suppression are common, potentially masking an infection. Although hypokalemia is more common in chronic hypothermia, hyperkalemia also occurs; the expected electrocardiographic changes can be obscured by hypothermia. Patients with renal insufficiency, metabolic acidoses, or rhabdomyolysis are at greatest risk for electrolyte disturbances.

Coagulopathies are common because cold inhibits the enzymatic reactions required for activation of the intrinsic cascade. In addition, thromboxane B_2 production by platelets is temperature dependent, and platelet function is impaired. The administration of platelets and fresh-frozen plasma is therefore not effective. The prothrombin or partial thromboplastin times or the international normalized ratio can be deceptively normal and contrast with the observed in vivo coagulopathy. This contradiction occurs because all coagulation tests are routinely performed at 37°C (99°F), and the enzymes are thus rewarmed.

REWARMING STRATEGIES

The key initial decision is whether to rewarm the patient passively or actively. *Passive external rewarming* simply involves covering and insulating the patient in a warm environment. With the head also covered, the rate of rewarming is usually $0.5^{\circ}-2^{\circ}$ C ($1.10^{\circ}-4.4^{\circ}$ F) per hour. This technique is ideal for previously healthy patients who develop acute, mild primary accidental hypothermia. The patient must have sufficient glycogen to support endogenous thermogenesis.

The application of heat directly to the extremities of patients with chronic severe hypothermia should be avoided because it can induce peripheral vasodilation and precipitate core temperature "afterdrop," a