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# Impact of Triiodothyronine on the **Survival of High-Risk Patients Undergoing Open Heart Surgery**

#### Abstract

Experimental and clinical studies have shown the beneficial effects of triiodothyronine (T<sub>3</sub>) following myocardial revascularization on cardiopulmonary bypass (CPB). In this study, open-label T<sub>3</sub> was administered to 68 high-risk patients undergoing open heart surgery. The New Jersey Risk Assessment was used to calculate the preoperative estimated surgical mortality. A loading dose of T<sub>3</sub> was administered: (a) at release of the aortic cross-clamp, (b) whenever the patient became CPB dependent, (c) if the patient exhibited low cardiac output after discontinuing CPB and (d) as pretreatment before initiating CPB. All therapeutic modalities were followed by a continuous T<sub>3</sub> infusion. Following T<sub>3</sub> therapy, CPB was discontinued in all patients. Based upon discriminant analysis, a total of 26 deaths were expected from the entire group, but only 7 patients died, therefore, the observed mortality was reduced by 72% (p < 0.007). The use of T<sub>3</sub> had a major impact on reducing surgical mortality, and may be advocated as a new therapeutic modality in patients with high estimated mortality undergoing open heart surgery.

# **Key Words**

Cardiopulmonary bypass Triiodothyronine

## Introduction

In cardiac surgery, the preoperative estimated mortality (EM) depends on both cardiac and noncardiac risk factors [1]. Relevant cardiac risks include: low left-ventricular ejection fraction, postmyocardial infarction new-onset unstable angina, congestive heart failure (CHF), re-do coronary surgery, combined valve and coronary artery surgery and others. Noncardiac risk factors such as diabetes, hypertension, chronic obstructive lung disease, and chronic renal failure, also have a direct impact on the surgical outcome (postoperative day 30) [2]. The EM can be calculated by using several models, such as the VA [1-3], STS [4, 5] and others. The New Jersey Risk Assessment was used in the current study [6].

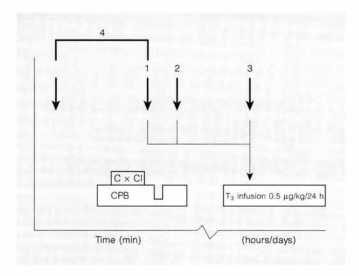
Despite advances in myocardial protection, myocardial depression often follows completion of open heart procedures [7-9]. There is a generalized body inflammatory response [10], complement activation, interleukin production and a profound derangement of the thyroid profile lasting for several days [11, 12]. The 'euthyroid sick syndrome' is characterized by a low FT<sub>3</sub>, increase in plasma reverse  $T_3$  (r $T_3$ ), normal or low total  $T_3$  (T $T_3$ ), thyroxine (T<sub>4</sub>), and thyroid-stimulating hormone (TSH) lev-

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**Fig. 1.** Modalities of  $T_3$  therapy: 1,  $T_3$  is administered at the time of release of the aortic cross clamp; 2,  $T_3$  is administered while patient remains CPB dependent; 3,  $T_3$  is administered in the post-CPB period; 4,  $T_3$  is administered before CPB (pretreatment) and a second dose at the time of release of the aortic cross clamp (C × Cl). Values are expressed in mean  $\pm$  SD.

els [13]. In sick patients, the lower the  $FT_3/rT_3$  ratios correlated with higher mortality [14, 15].

Experimental administration of T<sub>3</sub> in animals has shown a rapid reversal of the myocardial dysfunction which follows transient ischemic events, such as: (a) induction of experimental brain death [16-18], (b) stunned myocardium [19, 20], or (c) after prolonged cardioplegic arrest on cardiopulmonary bypass (CPB) [21, 22]. Similar observations were made in our preliminary clinical studies. After administering T<sub>3</sub> to 10 patients who were experiencing profound postcardiotomy myocardial depression, the T<sub>3</sub> enhanced the reversal of the myocardial dysfunction allowing for discontinuation of CPB, and also the use of intra-aortic balloon pump was no longer necessary after 1 h. This therapy also allowed for a rapid reduction in high inotropic requirements [23]. To confirm the beneficial effects of T<sub>3</sub>, we studied a group of low-risk (EM 1-5%) patients undergoing myocardial revascularization in a prospective, randomized, double-blind, placebo-controlled study was conducted, which confirmed the beneficial impact of T<sub>3</sub> therapy [24]. In the T<sub>3</sub>-treated patients, we observed significant hemodynamic improvement, requirement of less inotropic support and diuretics. The objective of this study was to observe the impact of  $T_3$ (in an open-label fashion) on high-risk patients undergoing open heart surgery.

#### **Patients and Methods**

From April 1992 to April 1994, at the University of South Florida affiliated hospitals (James A. Haley VA Hospital and The Tampa General Hospital), T<sub>3</sub> was administered to 68 patients undergoing open heart surgery. The mean age was 61.92 years (range 35-81 years), and there were 64 men and 4 women. There were 29 elective, 7 urgent and 32 emergent surgical procedures. Coronary artery bypass grafts were performed in 32 patients, valve procedures in 9 patients, combined coronary artery bypass grafts with valve procedures in 27 patients, and an orthotopic heart transplant in 1 patient. Eighteen patients required preoperative use of the intra-aortic balloon pump for control of post-myocradial infarction unstable angina, despite heparin and intravenous nitroglycerin, and 10 patients received inotropic support. At the completion of the surgical procedure, unexpectedly, 12 patients became CPB dependent, receiving T<sub>3</sub> following prolonged CPB support. The preoperative mean EM was 29.67% (range 3-58%), patients were informed of the EM and consent to administer T<sub>3</sub> was obtained prior to enrollment. In emergent conditions, or if the patient received sedation, consent was obtained from a legal representative. This protocol and informed consent forms were approved by the Institutional Review Board of the University of South Florida and the Research and Development Committee of the James A. Haley VA Hospital.

#### CPB Management: Perioperative Monitoring

Standard hemodynamic monitoring values were obtained from a Swan Ganz catheter, and an arterial line. Moderate hypothermia (28–32 °C) was used, and the mean arterial pressure was maintained in the range of 50–60 mm Hg with a flow of 2 liters/m² using a Sarns roller pump. Blood cardioplegia was administered in an antegrade and retrograde fashion. Epinephrine was the inotropic agent of choice; however, a combination of other inotropic drugs was used whenever indicated. Nitroglycerin and sodium nitroprusside were used to maintain systemic vascular resistance in the range of 600–800 dyn. CPB was discontinued progressively under various loading conditions, and inotropic requirements were adjusted accordingly.

In 12 patients, further CPB support was required for various intervals as the heart failed to sustain adequate hemodynamics, despite receiving epinephrine in excess of  $10 \,\mu\text{g/min}$  in combination with other inotropics (CI <1.8, MAP  $\leq$ 60 mm Hg and PCWP  $\geq$ 18 mm Hg). An attempt to discontinue CPB was initiated again. In the event of failure to support the circulation, the patients had a second, third or fourth bypass support runs.

## Plasma FT3 Measurement

Plasma FT<sub>3</sub> was measured by radioimmunoassay using a commercially available kit (Diagnostic Products Corporation, Calif., USA).

## T<sub>3</sub> Therapeutic Modalities

T<sub>3</sub> was administered using four different regimens (fig.1):

Regimen 1: n = 44,  $T_3$  loading bolus (2-3  $\mu$ g/kg) was administered at the time of release of the aortic cross clamp;

Regimen 2: n = 12,  $T_3$  (1–2  $\mu$ g/kg) was administered after the release of the aortic cross clamp, whenever the patient became CPB dependent.

Regimen 3: n = 4,  $T_3$  (1  $\mu g/kg$ ) was administered in the postoperative period to patients exhibiting a low cardiac output state, despite high IABP support;

Table 1. Risk stratification and expected mortality

#### A Percentage EM - preoperative

Groups		Preoperative EM1 according to discharge status, %						
		alive						
1	0-9	8, 4, 3, 9, 9	-					
2	10-19	14, 12, 16, 18, 14, 18, 17, 19, 12, 15, 11, 16	_					
3	20–29	18, 20, 25, 25, 22, 24, 20, 27, 24, 25, 29, 27, 27, 22, 20, 23, 24, 20, 25, 27, 27	21, 24					
4	30-39	33, 38, 36, 32, 35, 34, 30, 35, 37	32, 38					
5	40-49	41, 48, 49, 46, 48	45					
6	50-59	50, 57, 57, 55, 57, 51, 54, 54, 51	58, 56					

## **B** Observed mortality

Groups	n	Observed deaths	EM mean±SD, %	Expected deaths	O/E <sup>2</sup>	p value <sup>3</sup>
1	5	0	6.6±2.88	0.33	0	< 0.74
2 12 0			$15.16 \pm 2.62$	2.02	0	< 0.16
3	23	2	$24.00 \pm 2.72$	5.28	0.37	< 0.01
4	11	2	$34.54 \pm 2.62$	3.79	0.26	< 0.08
5	6 1		$46.16 \pm 2.92$	2.77	0.52	< 0.18
6	11	2	$54.54 \pm 2.80$	5.99	0.33	< 0.04
Total	68	7		20.17		p < 0.000

- Each number represents the preoperative estimated mortality for each patient.
- <sup>2</sup> O/E: Observed/Expected deaths; normally this ratio is equivalent to 1.
- 3 Two-tailed p values are expressed.

Regimen 4: n = 8,  $T_3$  pretreatment loading dose (1  $\mu$ g/kg) was administered 30–60 min before initiating CPB, followed by a second  $T_3$  bolus (1  $\mu$ g/kg) at the time of aortic cross clamp release.

Maintenance infusion: 0.5 μg/kg/24 h was administered from 1 to 3 days according to hemodynamic parameters.

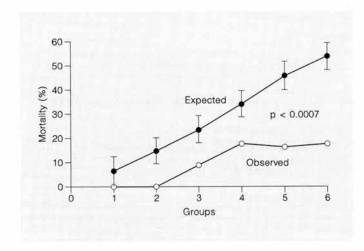
In 15 patients, following the loading bolus dose and during the infusion period,  $T_3$  kinetic studies were performed measuring the half-life of the distribution phase  $(t_{ij}\alpha)$ , half-life of the elimination phase  $(t_{ij}\beta)$  and the volume distribution.

## Statistical Analysis

Calculation of EM for each patient was carried out using the New Jersey Risk Stratification Model. All patients were stratified and allocated at 10% intervals according to the EM. The total expected deaths were calculated and the observed/expected ratio (O/E) expressed for each patient's interval. For the observed and estimated deaths, the level of significance was determined using a cumulative binomial probability distribution. The inotropic requirement for the CPB-dependent patients after the  $T_3$  bolus was compared using Student's t test. The values are expressed as mean  $\pm$  SD of the mean (table 1).

# Results

Of these 68 high-risk patients, CPB was discontinued in 56 patients, 12 remained CPB dependent. T3 was administered to 44 patients at the time of release of the aortic cross clamp, 4 patients received T<sub>3</sub> in the postoperative period. Eight had T<sub>3</sub> pretreatment followed by a second dose at the time of aortic cross clamp release. T<sub>3</sub> pretreatment was administered 30-60 min prior the initiation of CPB to patients experiencing low cardiac output state. While patients remained CPB dependent, T<sub>3</sub> was administered during the last CPB support run; a rapid cardiac functional recovery was observed, allowing the discontinuation of CPB support (table 2). Eight or more deaths were expected from the 12 CPB-dependent patients (including a primary cardiac graft failure following transplantation). Following the cardiac procedure, CPB was uneventfully discontinued in all patients. However, within 30 days, only 1 CPB-dependent patient died. Considering all patients together, 26 patients were expected to



die, however only 7 patients died. The EM was significantly reduced by 72% (p < 0.0007) (fig. 2). Of the 7 observed deaths (POD 30), 4 were cardiac; and 3 were noncardiac (table 3).

The preoperative FT<sub>3</sub> levels had a significant reduction from  $2.01 \pm 0.01$  to  $0.8 \pm 0.07$  pg/ml (p < 0.0001) prior to T<sub>3</sub> therapy. Following the initial T<sub>3</sub> bolus, FT<sub>3</sub> levels remained elevated for the first 2 h, however during

Fig. 2. Comparison of observed and expected mortality. 95% confidence intervals and the statistical differences are shown. Values are expressed in mean  $\pm$  SD.

Table 2. CPB-dependent patients

Age	Sex		EM PS C × CL CPB runs, min min 1 2 3		CPB runs, min					Cardiac status	Cardiac procedure	Inotropic support <sup>1</sup> Epi/µg/kg/min		Outcome
		%		3	4	5	pre T <sub>3</sub>	post T <sub>3</sub>						
69	М	14	U	114	186	30	20			AS, CAD, pre-op MI, IABP	AVR, CABG ×2	0.14	0.05	alive
44	M	10	Е	95	172	50	30			CAD, recent MI	CABG × 3	0.128	0.05	alive
62	М	18	Е	100	160	30	20	18		CAD, MR LVEF 25% dobut.	CABG × 3, MVR	0.266 norepi.	0.05 0	alive
50	M	14	Е	98	185	24				CAD, CABG, LVEF, 25%	redo CABG × 4	0.11	0.038	alive
66	M	24	E	66	239	48				CAD, CABG, recent MI, IABP	redo CABG × 3	0.10	0.025	alive
35	M	15	M	129	91	28	46	17	33	CAD, CM, HTx	oHTx norepi., Isup.	0.106 norepi., Isup.	0.053	alive
58	M	37	M	63	120	20				CAD, CRF, cath crash IABP	CABG × 4 norepi.	0.2 norepi., Amri	0.05 n	alive
74	M	24	Е	135	232	30				CAD, AS	CAGB × 1 AVR	0.22	0.22 POD7 RV MI	dead
62	M	7	Е	106	144	31				CAD, prev. MIs, LVEF 20%	VABG × 4	0.08	0.02	alive
54	M	46	Е	50	100	30				CAD, CABG MI, angina IABP	redo CABG × 2	0.15	0.05	alive
66	M	54	M	221	302	20				CAD, MR, MI angina, IABP LVEF 30%	CABG × 3, MVR	0.11	0.05	alive
53	М	16	M	103	120	26				CAD, acute MI post cath angina, IABP	CABG × 4	0.12	0.03	alive
Total	12			104.63 (13.79)	170.90 (18.16)	31.41 (2.55)	27.2 (5.10)	17.5 (0.5)	33			0.151 vs. (0.058)	0.06* (0.057)	11/12

EM = Estimated mortality; PS = preop. status; U = urgent; E = elective; M = emergent; A = aortic stenosis;

C × CL = aortic cross clamp; Epi = epinephrine, oHTx = Orthotopic heart transplant; IABP = intra-aortic balloon pump.

Epinephrine requirement was reduced from 0.15 (0.058) to 0.06 (0.057)  $\mu$ g/kg/min p < 0.0001.

the infusion interval the FT<sub>3</sub> levels were within the normal range. The  $t_{\nu_2}\alpha$  was of 0.42  $\pm$  0.03 h and the  $t_{\nu_2}\beta$  was of 1.69  $\pm$  0.08 h. The volume distribution was 41.66 liters/kg.

## Discussion

In this open study, the beneficial effects of T<sub>3</sub> following open heart surgery were confirmed. There was a significant improvement on the survival of high-risk patients undergoing a variety of cardiac procedures. The beneficial effects of T<sub>3</sub> were seen not only in patients undergoing valvular surgery with normal coronary arteries, but also in patients undergoing myocardial revascularization in the presence of acute MI and new-onset angina exhibiting severaly compromised left-ventricular function and preoperative 'stunning'.

The beneficial effects of T<sub>3</sub> are multifactorial, impacting not only on the cardiovascular system but on other organs as well [25]. At the cardiac level it is well known

that T<sub>3</sub> affects various ATPase [26, 27] and also upregulation of beta-receptors [27, 28]. It also increases coupling of the beta-receptor to adenylate cyclase [29]. T<sub>3</sub> also plays a role in calcium uptake and calcium release from the sarcoplasmic reticulum into the cytosol and activates various calcium-dependent ATPases [30-32], thus rapidly mobilizing and recompartmentalizing free ionic calcium from the cytosol into the sarcoplasmic reticulum [33, 34]. If this is the case, and also occurs in other organs, the calciuminduced injury which resembles the oxygen free radical injury [35] occurring during CPB may be reduced. T<sub>3</sub> also enhances metabolism at the mitochondrial level [36, 37] by increasing the activity of the adenine nucleotide translocase facilitating ATP mobilization from the mitochondria into the cytosol, thus, yielding high-energy phosphates required for the myosin ATPase [38].

The observed inotropic effects following  $T_3$  administration have been documented within several minutes, and are probably extra-nuclear related, bypassing the DNA-RNA protein synthesis and independent of  $\beta$ -receptor stimulation. This rapid response is of great relevance

Table 3. Surgical mortality at postoperative day (POD) 30

Patient	Age	Sex	EM %	Status	Pre-diagnosis	Surgical procedure	Cause of death
1	72	М	58	M	MI, new onset angina, MTG, hep, IABP	CABG × 3	POD 20, SP initial course, transferred to floor, acute abdomen, laparotomy POD 18, small bowel infarct
2	74	M	24	Е	CAD, AS, MI	AVR, CABG × 1 LAD endarterectomy	POD 7, new perioperative MI, ARDS, inotropes, low cardiac output
3	73	M	32	Е	CAD, porcelain aorta	CABG × 3	POD 3, multiple perioperative MIs possibly from aortic emboli
4	62	M	21	M	CAD, unstable angina, hep, MTG, LVEF 20%	CABG × 3	POD 18, uneventful ICU course mediastinitis; POD 15, cardiac-arrhythmia asystole
5	60	M	56	M	CABG, MI, PTCA, acute graft dissection	redo CABG × 4	POD 9, perioperative MI, low cardiac output
6	74	M	45	Е	aortic prost. valve stenosis MR +4, TR +4	redo AVR, MVR, Tann	POD 5, multiorgan failure, sepsis, PM severe cirrhosis
7	791	M	38	Е	AS, MR, CAD	AVR (SJ No. 19) M. Ann CABG × 2	POD 29, extubated, then ARDS pulmonary edema; severe aortic valve gradient ± 90 mm/Hg mediastinitis, sepsis

Patient 7 had a number 19 aortic prosthesis, postoperative gradient by TEE measured 80-110 mm Hg. The LV and RV functions remained good.

Redo AVR was not performed as the patient developed ARDS and sepsis (mediastinitis).

to cardiac surgery [39–41]. This rapid recovery was also observed after inducing myocardial stunning in dogs [20]. By using load-independent pressure-volume relationship and sonomicrometers  $T_3$  allowed total recovery of the myocardial dysfunction.

In our study, T<sub>3</sub> pretreatment was used for the first time in patients prior to the initiation of the open heart procedure. The rationale for this therapeutic modality was based on the previous experience of administering T<sub>3</sub> to brain-dead animals and unstable human brain-dead organ donors depending on high inotropic support [42–44]. The impact of T<sub>3</sub> resulted in a significant hemodynamic and biochemical improvement allowing rapid inotropic reduction. Following cardiac transplantation, these marginal donor hearts performed well in recipients [45].

Eight patients in a study exhibited a marked reduction of the systemic vascular resistance requiring the administration of norepinephrine in order to maintain adequate tissue perfusion pressure; however, diuresis remained excellent and no deleterious effects were observed, thus reconfirming that  $T_3$  at the vascular level is a potent vasodilator [46]. It is quite possible that the  $T_3$  loading dose of  $3 \mu g/kg$  is the optimal pharmacological dose as is denoted by the marked vasodilation which followed administration. The concept that no thyroid replacement should be administered to patients exhibiting the 'euthyroid sick

syndrome' as in those exposed to acute insults [47–49] seems no longer to be invalid, particularly in patients undergoing open heart surgery on CPB.

The marked reduction of half-life of the  $\beta$ -phase following exogenous  $T_3$  administration was found to be approximately 8% of the endogenously produced  $T_3$ . This rapid reuptake may be directly related to the tissue  $T_3$  depletion following CPB.

The conclusions of this study are limited by its openlabel design and lack a placebo control. If, however, our assumption proves to be correct, T<sub>3</sub> may improve the survival of high-risk patients undergoing cardiac surgery.

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#### References

- 1 Grover F, Johnson R, Marshall G, Hammermeister KE: Factors predictive of operative mortality among coronary artery bypass subsets. Ann Thorac Surg 1993;56:1–11.
- 2 Hammermeister KE, Burchfiel C, Johnson R, Grover FL: Identification of patients at greatest risk for developing major complications at cardiac surgery. Circulation 1990;82:380–389.
- 3 Grover FL, Hammermeister KE, Burchfiel C, Cardiac Surgeons of the Dept. of Veterans Affairs: Initial report of the veterans administration preoperative risk assessment study for cardiac surgery. Ann Thorac Surg 1990;50:12–28.
- 4 Clark RE: The Society of Thoracic Surgeons National Data Base Status Report. Ann Thorac Surg 1993;57:27–32.
- 5 Edwards ED, Clark RE, Schwartz M: Coronary artery bypass grafting: The Society of Thoracic Surgeons National Data Base Experience. Ann Thorac Surg 1994;57:20–26.
- 6 Parsonnet V, Dean D, Burstein AD: A method of uniform stratification of risk for evaluating the results of surgery in acquired adult heart disease. Circulation 1988(suppl 1);79:3–12.

- 7 Hearse DJ: Stunning: A radical review. Cardiovasc Drugs Ther 1991;5:853–876.
- 8 Keith A, Fox A, Bergmann SR, Sobet BE: Pathophysiology of myocardial reperfusion. Annu Rev Med 1985;36:125-144.
- 9 Kukielka GL, Hawkins KH, Michael L, Manning AM, Youker K, Lane C, Entman ML, Smith CW, Anderson DC: Regulation of intercellular adhesion molecule-1 (ICAMO1) in ischemic and reperfused canine myocardium. J Clin Invest 1993;92:1504–1516.
- 10 Edmunds LH: Systemic inflammatory responses secondary to cardiopulmonary bypass. Am J Surg 1993(CME suppl):4–9.
- 11 Robuschi G, Medici D, Fesani F, Barboso G, Montermini M, d'Amato L, Gardini E, Borciani E, Dall'Aglio E, Salvi M: Cardiopulmonary bypass: A low T<sub>4</sub> and T<sub>3</sub> syndrome with blunted thyrotropin (TSH) response to thyrotropin-releasing hormone (TRH). Horm Res 1986:23:151-158.
- 12 Clark RE: Thyroid and other endocrine responses to cardiopulmonary bypass. Am J Surg 1993(CME suppl):10–14.

- 13 Wartofsky L, Burman KD: Alterations in thyroid function in patients with systemic illness: The 'euthyroid sick syndrome'. Endocr Rev 1982;3:164–217.
- 14 Slag MF, Morley JE, Elson MK: Hypothyroxinemia in critically ill patients as a predictor of high mortality. JAMA 1981;245:43–45.
- 15 Hamilton MA: Prevalence and clinical implications of abnormal thyroid hormone metabolism in advanced heart failure. Ann Thorac Surg 1993;56:548–553.
- 16 Novitzky D, Wicomb WN, Cooper DKC, Frazer R, Barnard CN: Electrocardiographic hemodynamic and endocrine changes occurring during experimental brain death in the Chacma baboon. Heart Transplant 1984;4:63.
- 17 Novitzky D, Copper D, Reichart B: Hemodynamic and metabolic responses to hormonal therapy in brain-dead potential organ donors. Transplantation 1987;43:852–854.
- 18 Novitzky D, Cooper D, Chaffin J, Greer AE, DeBault LE, Zuhdi N: Improved cardiac allograft function following triiodothyronine therapy to both donor and recipient. Transplantation 1990;49:311–316.

- 19 Novitzky D, Matthews N, Shawley D, Cooper DKC, Zuhdi N: Triiodothyronine in the recovery of stunned myocardium in dogs. Ann Thorac Surg 1991;51:10-17.
- 20 Yokoyama Y, Novitzky D, Deal MT, Snow TR: Facilitated recovery of cardiac performance by triiodothyronine following a transient ischemic insult. Cardiology 1992;81:34-
- 21 Novitzky D, Human P, Cooper D: Inotropic effect of triiodothyronine following myocardial ischemia and cardiopulmonary bypass: An experimental study in pigs. Ann Thorac Surg 1988-45-50-55
- 22 Novitzky D, Human P, Cooper D: Effect of triiodothyronine (T3) on myocardial high energy phosphates and lactate after ischemia and cardiopulmonary bypass. J Thorac Cardiovasc Surg 1988;96:600-607.
- 23 Novitzky D, Cooper DKC, Swanepoel A: Inotropic effect of triiodothyronine (T3) in low cardiac output following cardioplegic arrest and cardiopulmonary bypass: Initial experience in patients undergoing open heart surgery. Eur J Cardiothorac Surg 1989;3:140-145.
- 24 Novitzky D, Cooper D, Barton I, Greer A. Chaffin J, Grim J, Zuhdi N: Triiodothyronine as an inotropic agent after open heart surgery. J Thorac Cardiovasc Surg 1989;98:972-978.
- 25 Pienaar H, Schwartz I, Roncone A, Lotz Z, Hickman R: Function of kidney grafts from brain-dead donor pigs. The influence of dopamine and triiodothyronine. Transplantation 1990:50:580-582.
- 26 Rudinger A, Mylotte K, Davis P, Davis F, Blas S: Rabbit myocardial membrane Ca2+ adenosine triphosphatase activity: Stimulation in vitro by thyroid hormone. Biochem Biophys 1984;229:379-385.
- 27 Davis PJ: Cellular actions of thyroid hormones; in Braverman LE, Utinger RD (eds): Werner and Ingbar's the Thyroid, ed 6. Philadelphia, 1991.

- 28 Ginsbert AM, Clutter WE, Shah SD, Cryer PE: Triiodothyronine-induced thyrotoxicosis increases mononuclear leukocyte beta-adrenergic receptor density in man, J Clin Invest 1981;67: 1785-1791.
- 29 Segal J: Action of the thyroid hormone at the level of the plasma membrane. Endocr Res 1989:15:619-649
- Rudinger A, Mylotte KM, Davis PJ, Davis FB. Blas SD: Rabbit myocardial membrane Ca++ ATPase activity: Stimulation in vitro by thyroid hormone. Arch Biochem Biophys 1984; 229:379-385.
- 31 Segal J: In vivo effect 3,5,3'-triiodothyronine on calcium uptake in several tissues in the rat: Evidence for a physiologic role for calcium as the first messenger for the prompt action of thyroid hormone at the level of the plasma membrane. Endocrinology 1990;127:17-24.
- Kim D, Smith T: Effects of thyroid hormone on calcium handling in cultured chick ventricular cells. J Physiol (Lond) 1985;364:131-149.
- 33 Warnick PR, Davis PJ, Davis FB: Rabbit skeletal muscle sarcoplasmic reticulum Ca2+-ATPase activity: Stimulation in vitro by thyroid hormone analogues and bipyridines. Biochim Biophys Acta 1993;115:184-190.
- Toyofuku T, Kurzydlowski K, Tada M, MacLennan DH: Identification of regions in the Ca2+-ATPase of sarcoplasmic reticulum that affect functional association with phospholamban. J Biol Chem 1993;268(4):2809-2815
- 35 Krause SM, Jacobus WE, Becker LC: Alterations in cardiac sarcoplasmic reticulum calcium transport in the postischemic 'stunned' myocardium. Circ Res 1989;65:526-530.
- 36 Sterling K: Direct thyroid hormone activation of mitochondria: The role of adenine nucleotide translocase. Endocrinology 1986;119:292-295.
- 37 Davis PJ: Cellular actions of thyroid hormones; in Braverman LE, Utinger RD (eds): Warner and Ingbar's the Thyroid, ed 6. Philadelphia, 1991.
- 38 Hoh JFY, McGrath PA, Hale PT: Electrophoretic analysis of multiple forms of cardiac myosin: Effect of hypophysectomy and thyroxine replacement. J Mol Cell Cardiol 1978;10: 1053-1076.

- 39 Snow TR, Deal MT, Connelly TS, Yokoyama Y, Novitzky D: Acute inotropic response to rabbit papillary muscle to triiodothyronine. Cardiology 1992;80:112-117.
- 40 Dyke CM, Yeh T, Lehman JD, Abd-Elpattah A, Ding M, Wechsler AS: Triiodothyronineenhanced left ventricular function after ischemia injury. Ann Thorac Surg 1991;52:14-19.
- 41 Dyke CM, Ding M, Abd-Elfattah AS, Loesser K, Dignan RJ, Wechsler AS, Salter DR: Effects of triiodothyronine supplementation after myocardial ischemia. Ann Thorac Surg 1993; 56:215-222.
- 42 Taniguchi S, Kitamuras S, Kawachi K, Doi Y, Aoyama N: Effects of hormonal supplements on the maintenance of cardiac function in potential donor patients after cerebral death. Eur J Cardiothorac Surg 1992;6:96.
- 43 Orlowski JP, Spees EK: Improved cardiac transplant survival with thyroxine treatment of hemodynamically unstable donors: 95.2% graft survival at 6 and 30 months. Transplant Proc 1993;25:1535.
- Novitzky D, Wicomb WN, Cooper DKC, Tjaalgard MA: Improved cardiac function following hormonal therapy in brain-dead pigs: Relevance to organ donation. Cryobiology 1987;24:1-10.
- 45 Novitzky D, Cooper DKC, Chaffin JS, Greer AE, Debault LE, Zuhdi N: Improved cardiac allograft function following triiodothyronine (T<sub>3</sub>) therapy to both donor and recipient. Transplantation 1990;49:311-316.
- Klein I: Thyroid hormone and the cardiovascular system. Am J Med 1990;88:631-637.
- Dulchavsky SA, Lucas CE, Ledgewood AM, Grabow D: Triiodothyronine (T<sub>3</sub>) improves CV function during hemorrhagic shock. Circ Shock 1993:39:67-73.
- Klein I: Treatment of myxedema-associated cardiogenic shock. Ann Intern Med 1993;119: 168-169
- 49 Hesch RD, Husch M, Kodding R, Hoffken B, Mayer T: Treatment of dopamine-dependent shock with T3. Endocr Res Commun 1981;8: 229-237.

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