

Dilated Cardiomyopathy and Hypothyroidism with concomitant CAD -a debatable scenario

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Abstract

The concept of dilated cardiomyopathy with concomitant coronary artery disease and hypothyroidism is yet a matter of debate although many theories have been postulated with this regard. We report overt heart failure due to dilated cardiomyopathy in an elderly lady with significant coronary artery disease and hypothyroidism. Early suspicion, evaluation and judicious use of thyroxine with appropriate anti ischaemic measures proved beneficial with a better prognostic outcome in this patient.

Index terms— heart failure, dilated cardiomyopathy, ischemic heart disease, hypothyroidism. CAD coronary artery disease.

1 INTRODUCTION

: heart failure, dilated cardiomyopathy, ischemic heart disease, hypothyroidism. CAD-coronary artery disease. Thyroid hormone has many effects on the heart and vascular system. [1] The heart is very sensitive to alterations in serum thyroid levels. Many of the clinical manifestations of hyperthyroidism and hypothyroidism are due to the ability of thyroid hormone to alter cardiovascular hemodynamics. [2] Thyroid hormone metabolism is altered in many patients with acute or chronic cardiac disease, as it is in patients with other non-thyroidal illnesses. Cardiac manifestations of thyroid hormones are due to dyslipidemia, accelerated atherogenesis, reduced heart rate, contractile states of myocardium and pericardial effusion.

The prevalence and clinical importance of myocardial dysfunction in hypothyroidism are generally overlooked. Nonspecific histological abnormalities have been demonstrated repeatedly in the hearts of myxoedema patients since first reported in 1888 in report of a committee of the Chemical Society of London. [3] The structural changes together with haemodynamic changes in heart of a hypothyroid patient termed as hypothyroid cardiomyopathy [4] [5] [6] has shown a good response to thyroxine replacement.

Ischaemic cardiac events have also been implicated in causing transient thyroid dysfunction. But whether the cardiomyopathy associated with both ischemic heart disease and hypothyroidism are interrelated is still a matter of debate as significant improvement has been seen in patients treated concurrently for the two different conditions.

2 II.

3 CASE REPORT

A 61 year old lady presented to the emergency department of a tertiary institution with exertional dyspnoea since 1 week with no documented medical history. On examination the patient was mildly cyanosed, pulse rate 120 beats per minute; regular rhythm, blood pressure 110/90 mmHg, respiratory rate 32 cycles/min, saturation of O₂ 85% with elevated JVP. Cardiovascular examination showed tachycardia with gallop rhythm and bilateral basal crepitations. A clinical diagnosis of heart failure was made. All preliminary investigations were within normal range except the lipid profile which was altered with total cholesterol-320 mg/dl, LDL cholesterol -180 mg/dl, HDL-40 mg/dl, TG -380 mg/dl with normal values of CPK and Troponin-I. Electrocardiogram showed

5 DISCUSSION

43 sinus tachycardia with no ST -T changes. Chest x-ray showed cardiomegaly (figure ?? a) with pruning of
44 upper lobar veins and peri hilar congestion. 2D Echocardiography showed global hypokinesia with an inter
45 ventricular septal thickness of 8.1 mm, mild mitral regurgitation, no regional wall motion abnormalities, minimal
46 pericardial effusion and an left ventricular ejection fraction (LVEF) of 27%. She was decongested with diuretics
47 and recovered symptomatically. An emergency coronary angiography was performed, which revealed a triple
48 vessel disease with blocks of -left anterior descending (LAD) 60 %, left circumflex (LCX) 100%, mid right
49 coronary artery (RCA) 100%. Respecting her LVEF of 27% she was subjected to conventional treatment. A
50 true benefit of an interventional revascularization in this patient was a dilemma at this point, hence this patient
51 was contemplated for radio nucleotide 99m technetium (Tc) -resting myocardial perfusion study (figure ??,2)
52 which showed severe degree of resting myocardial perfusion defects in the anterior wall, inferior and lateral walls
53 including the apex and septum, corresponding to LAD, RCA, LCX territories and minimal to moderate degree of
54 viable myocardium, LV dilatation with evidence of systolic and diastolic dysfunction .Her above follow up pointed
55 out to a ischemic cause of the underlying heart failure and she was promptly started on conventional oral anti
56 ischemic measures(Aspirin 150mg, clopidogrel 75mg), ACE inhibitors(ramipril 1.25mg) ,statins (atorvastatin
57 40mg) , aldosterone antagonist (T Author ? ? : Professor, Post graduate student, Al Ameen medical college
58 and Hospital , Bijapur-586108, Karnataka, India. eplerenone 25mg) and low dose diuretics.(torsemide 20 mg).
59 With this therapy her symptoms were brought under control. In the mean time her thyroid profile was awaited
60 which pointed towards frank hypothyroidism. She was started cautiously with Levothyroxine 0.25 micro gram
61 per day which was gradually built upto a dose 0.1 mg over 3 weeks duration. Following this treatment she showed
62 significant improvement in her symptoms and was later discharged after fixing the dose of thyroxine at 0.1mg/
63 day.

64 During serial follow ups, 4 months after her discharge, she was asymptomatic, active and able to carry out her
65 routine activities .She was reinvestigated to study her present status and therapeutic response. Her chest x-rays
66 (figure ?? b, c) taken then showed significant reduction in cardiac size. Echocardiography showed improved
67 LVEF to 55%. Her lipid profile and thyroid function test were also within normal ranges. She was continued on
68 her medications and has been doing well till date.

69 4 III.

70 5 DISCUSSION

71 Thyroid hormone has many effects on the heart and vascular system.* Many of the clinical manifestations
72 of hyperthyroidism and hypothyroidism are due to the ability of thyroid hormone to alter cardiovascular
73 hemodynamics.The hemodynamic effects of hypothyroidism are opposite to those of hyperthyroidism, although
74 the clinical manifestations are less obvious. Prompt evaluation here made the diagnosis of congestive
75 cardiomyopathy. We believed this to be related to the underlying coronary pathology with respect to her
76 altered lipids, low ejection fraction, poor LV systolic function and global hypokinesia supported with coronary
77 angiogram. Radionuclide tech 99 resting myocardial perfusion scan revealed significant perfusion defects with
78 viable myocardium. PET studies of O2 consumption in patients with hypothyroidism have revealed that
79 myocardial work efficiency is lower than in normal subjects.[7]Significant dyslipidemia in a slim elderly patient
80 (BMI-20) prompted us to investigate her thyroid status as such an accelerated coronary atherosclerosis due
81 to hypercholesterolemia in hypothyroidism and post menopausal ladies has already been postulated* Some
82 theories have explained overt hypothyroidism to occur following an acute coronary event or acute myocardial
83 infarction,*but the phenomena is a sub clinical state of hypothyroidism and in heart failure, patients have low
84 serum T3 concentration and the degree is proportional to severity of heart failure as per NYHA functional
85 classification. ??8] We were in a dilemma as to whether the heart failure has depressed the thyroid hormones
86 or hypothyroidism per se is only the true cause for this cardiomyopathy. Here significant elevation of TSH more
87 than 150 and significant reduction in T3 and T4 made the diagnosis of hypothyroid cardiomyopathy. We had
88 initially thought of IV T3 as an immediate therapy to tide over this crisis but due to its non availability we treated
89 this patient cautiously with thyroxine initiating with the lowest possible dose, gradually building up the dose
90 to a maximum of 0.1 mg within 6 weeks. In fact risk versus benefits with thyroxine therapy in elderly patients
91 with concomitant coronary artery disease were thought seriously as thyroxine is known to improve the cardiac
92 contractility and reduce the peripheral vascular resistance and has no effect in improving the LVEF .Theories
93 have explained maximum beneficial effects of thyroxine in patients who were diagnosed to have heart disease in
94 long standing hypothyroidism*but in our case it was a risk as patient was tachycardic. Many of the patients with
95 severe heart failure in hypothyroidism with significantly compromised LVEF, poor LV systolic function and a
96 jeopardized myocardium are expected to have prolonged QT interval and abrupt initiation of thyroxine therapy
97 in them may culminate with torse de pointes, ventricular arrhythmias and a premature sudden cardiac death.

98 Whether the thyroid condition in this case was a separate preexisting entity precipitating the underlying
99 cardiac events or whether it was precipitated by the cardiac event was yet to be explained.

100 **6 IV.**

101 **7 CONCLUSION**

102 Patients with thyroid diseases often have symptoms and signs indicating changes in cardiovascular hemodynamics.
103 Indeed, symptoms and signs referable to the cardiovascular system may be the only manifestations of thyroid
104 dysfunction, and thyroid function should therefore be assessed by the measurement of serum thyrotropin
105 concentrations in patients with cardiovascular disease. Patients with cardiovascular disease, like patients with
106 other nonthyroidal illnesses, have changes in thyroid hormone metabolism that may alter cardiac function.
107 Although some data suggest that the thyroid replacement therapy may benefit some patients with cardiovascular
disease, further studies are required to establish specific treatment recommendations. ^{1 2 3}



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Figure 1: Figure 1 Figure 2 DilatedFigure 3 :

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