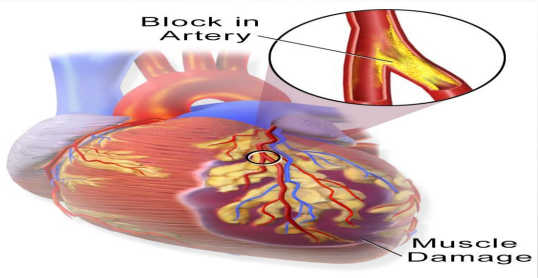
CARDIOVASCULAR SYSTEM BIOCHEMICAL MARKERS FOR MI

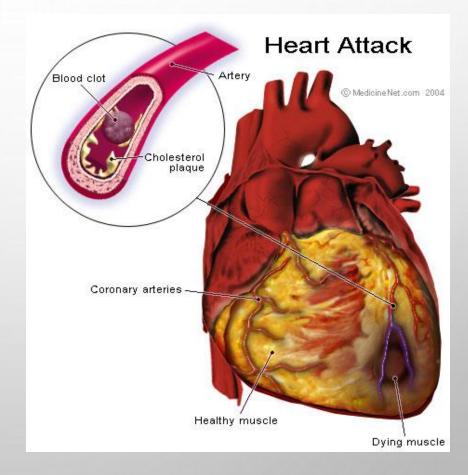


Heart Attack

DR. HEBA M. ABD EL KAREEM ASSISTANT PROFESSOR OF MEDICAL BIOCHEMISTRY AND MOLECULAR BIOLOGY

Acute Myocardial Infarction

- An imbalance between the supply of oxygen and the myocardial demand resulting in myocardial ischemia.
- A rapid development of myocardial necrosis caused by prolonged ischemia resulting in an irreversible myocardial injury.
- The development of infarction or ischemia will depend on the degree of occlusion or the presence of collateral blood flow.

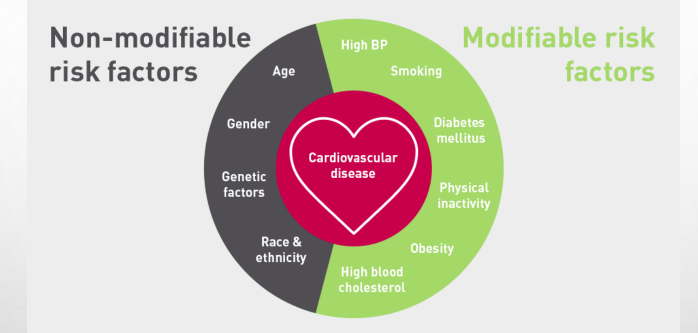


Biochemical Changes

ischemia to myocardial muscles (with low O_2 supply) anaerobic glycolysis increased accumulation of Lactate decrease in pH activate lysosomal enzymes disintegration of myocardial proteins cell death & necrosis

clinical manifestations (chest pain) BIOCHEMICAL MARKERS changes release of intracellular contents to blood

RISK FACTORS



- LDL-C is most important **atherogenic** particle.
- Apo B: Only apoprotein on LDL. ApoA1 is often used as a biomarker for prediction of CVD.
- ApoB100 / ApoA1 ratio is more effective at predicting heart attack risk, in patients who had had an acute MI, than either the ApoB100 or ApoA1 measure alone.

MYOCARDIAL INFARCTION

- Many patients with myocardial infarction have a typical history of crushing <u>central chest pain</u>, perhaps radiating to the arm or jaw, associated with typical ECG changes.
- myocardial infarction can, however, present <u>atypically</u>, or even be clinically silent, particularly in the elderly.
- The clinical evaluation often is limited by atypical symptoms, in most patients the initial ECG is non-diagnostic.
- The role of cardiac markers in the diagnosis and treatment of patients with chest pain and suspected AMI has evolved considerably.

WHAT ARE THE INVESTIGATIONS?

- ECG. chest x-ray coronary angiogram
- Lipid profile

- serum cardiac enzymes & proteins.

WHO Diagnosis of Acute Myocardial Infarction (AMI)

Presence of two of the three criteria:

- 1. History of characteristic chest pain.
- 2. Electrocardiographic changes.
- 3. Typical **pattern of serum cardiac enzyme & proteins** rise, peak and return to reference range.
 - However, in 1999, European Society of Cardiology and the American College of Cardiology
 - Sensitive biomarkers for the diagnosis of AMI
 - Cardiac troponins (cTn) is the gold standard.

IDEAL CARDIAC MARKER CHARACTERISTICS

- Cardiac specific. specific to myocardial muscle cells (no false positive).
- Sensitive: can detect minor damage. no miss of positive cases (no false negative)
- **Prognostic**: relation between plasma level & extent of damage
- Rises **soon** after plaque rupture.
- Elevated over a sustained period of time.
- Easy to measure, fast assay.
- Diagnostic utility verified by clinical studies.

QUESTIONS ANSWERED BY MARKERS OF CARDIAC DAMAGE

- RULE IN/OUT AN ACUTE MI
- CONFIRM AN OLD MI (SEVERAL DAYS)
- MONITOR RE-INFARCTION
- MONITOR THE SUCCESS OF THROMBOLYSIS

Biochemical markers in myocardial ischemia /necrosis

1. Cardiac Enzymes (isoenzymes):

- Total CK, CK-MB activity, CK-MB mass
- Aspartate aminotransferase (AST), Lactate dehydrogenase (LDH),.
- Glycogen phosphorylase BB (GPBB).

2. Cardiac proteins:

- Myoglobin & Troponins
- Ischemia Modified Albumin
- Heart-Fatty Acid binding protein (H-FABP).
- 3. Micro RNA (miRNA)

BIOCHEMICAL MARKERS IN MYOCARDIAL ISCHAEMIA /NECROSIS

OBSOLETE

- ASPARTATE AMINOTRANSFERASE -TOTAL CK - LACTATE DEHYDROGENASE

ESTABLISHED

- TROPONINT TROPONINI CK/MB MYOGLOBIN
- **EMERGING**
- MICRO RNA (MIRNA)
- HEART FATTY ACID-BINDING PROTEIN (H-FABP)
- ISCHEMIA-MODIFIED ALBUMIN
- GLYCOGEN PHOSPHORYLASE BB (GPBB)
- COPEPTIN B-TYPE NATRIURETIC PEPTIDE
- GROWTH DIFFERENTIATION FACTOR 15 PREGNANCY-ASSOCIATED PLASMA PROTEIN A

LABORATORY INVESTIGATIONS

SPECIMEN COLLECTION:

- **SERUM** IS THE SPECIMEN OF CHOICE
- HEPARINIZED PLASMA IS ACCEPTABLE
- **VENOUS WHOLE BLOOD** FOR RAPID CARDIAC TROPONIN T.
- SALIVA

COLLECTION TIME:

- SERIAL SPECIMENS COLLECTED AT APPROPRIATE TIME INTERVALS.
- SERIAL MEASUREMENTS ARE MOST USEFUL
- SAMPLES ARE DRAWN ON ADMISSION

AT 2-4 HOURS

AT 6-8 HOURS

AT 12 HOURS

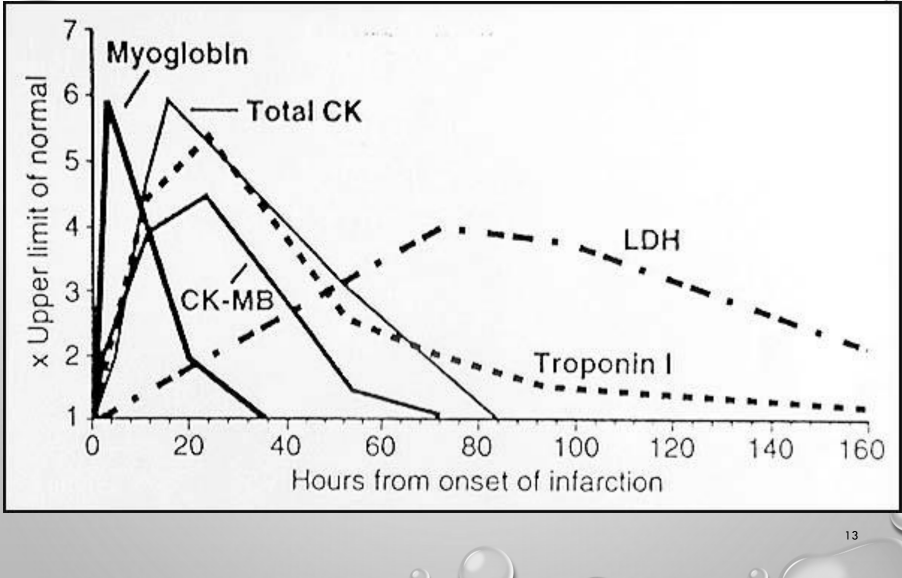
MYOGLOBIN

- O2-binding protein (heme-containing protein).
- Released from skeletal and heart muscle when damaged.
- Rapidly cleared by kidneys (not long term marker).
- Its level varies with gender, age, physical activity.
- More sensitive than CK, CK-MB activities.
- myoglobin is <u>not cardiac specific</u>, better used in conjunction with other markers. Increased in patients with skeletal muscle disease and chronic renal failure

MYOGLOBIN

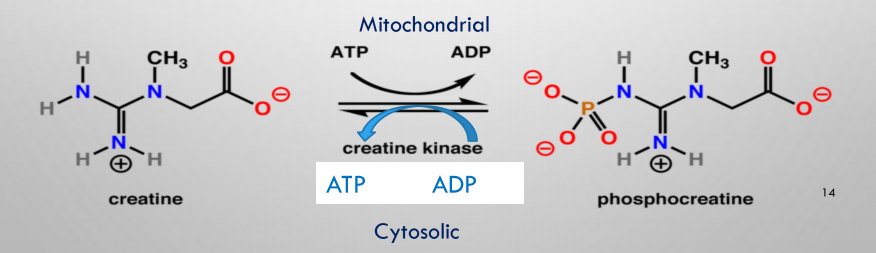
- IT **STARTS** TO RISE WITHIN **1-4 H**
- DETECTED BETWEEN 6-9 H IN NEARLY ALL AMI PATIENTS FROM CHEST PAIN.
- **<u>RETURNS</u>** TO BASE LINE LEVELS WITHIN 18-24 H.
- IF MYOGLOBIN ARE NORMAL 8H AFTER PAIN AMI CAN BE <u>RULES OUT</u>.
- [CK-MB IS PREFERRED THAN MYOGLOBIN IN PATIENTS WHO ARE ADMITTED LATER THAN **10-12 H** AFTER PAIN].

Biochemical markers of MI



CREATINE KINASE (CK)

- Creatine kinase acts as a regulator of high-energy phosphate production and utilization within contractile tissues.
- Cytoplasmic CK is a dimer, composed of M and/or B subunits, which associate forming CK-MM, CK-MB and CK-BB isoenzymes
- CK catalyses the conversion of creatine and consumes ATP to create phosphocreatine (PCr) and ADP.
- This CK enzyme reaction is <u>reversible</u>, such that also ATP can be generated from PCr and ADP.



CREATINE KINASE (CK)

- **CK-MM** is the main isoenzyme found in skeletal >> Cardiac muscles.
- <u>CK-MB</u> is found mainly in cardiac muscle Trace amounts of CK-MB are found in skeletal muscle.
- <u>CK-BB</u> is the predominant isoenzyme found in brain, colon, ileum, stomach and urinary bladder.

CK- TOTAL

- A RAISED PLASMA TOTAL CK ACTIVITY, DUE TO ENTIRELY <u>CK-MM</u> MAY FOLLOW:
- > SKELETAL MUSCLE DISEASE.
- > RECENT INTRAMUSCULAR INJECTION
- > EXERCISE
- > SURGERY.
- (NON SPECIFIC)
- LIMITED PROGNOSTIC VALUE.

CK-MB

- High specificity. more specific than total CK BUT: less specific than troponin I.
- Gold standard as cardiac marker (was).
- It takes at least 4-6 h to increase.
- Peak levels at 12-24 h.
- Return 2-3 days.
- useful for early diagnosis of MI
- useful for diagnosis re-infarction

• <u>CK-MB (MASS)</u>

MASS ESTIMATION <u>BETTER</u> THAN ACTIVITY. TO INCREASE SPECIFICITY, RATIO (RELATIVE INDEX) RELATIVE INDEX = CK-MB MASS / CK ACTIVITY.

CK-MB (MASS)

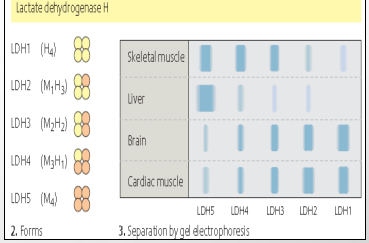
- If ratio >>> 3 indicative of AMI rather than skeletal muscle damage.
- CK/MB isoenzyme is not myocardium-specific occurring for instance in a small amount in skeletal muscle.
- Its use in the diagnosis AMI is considered acceptable only in cases where cTn assays are unavailable.
- The one advantage of CK-MB over the troponins is the early clearance that helps in the detection of re-infarction.

ASPARTATE TRANSAMINASE (AST)

- <u>HEPATIC CONGESTION</u> DUE TO RIGHT-SIDED HEART DYSFUNCTION MAY CONTRIBUTE TO THE RISE OF PLASMA AST ACTIVITY. A <u>NON-SPECIFIC</u> MARKER OF MI
- IF THERE IS <u>PRIMARY HEPATIC DYSFUNCTION</u>, PLASMA AST RISES WHEREAS LDH1 ACTIVITY USUALLY REMAINS NORMAL.
- THE SEQUENCE OF CHANGES IN <u>PLASMA AST ACTIVITY IN MI</u> IS <u>SIMILAR</u> TO THOSE OF <u>CK</u>.
- AST AND LDH MEASUREMENTS <u>ARE RARELY OF PRACTICAL VALUE</u> IN THE MANAGEMENT OF PATIENTS WITH SUSPECTED MYOCARDIAL INFARCTION.
- EXCEPTIONALLY, WHEN A PATIENT WITH CHEST PAIN PRESENTS LATE, MEASUREMENT OF LDH MAY BE HELPFUL AS THIS ENZYME REMAINS
 ELEVATED IN THE PLASMA FOR SEVERAL DAYS FOLLOWING MYOCARDIAL INFARCTION.

Lactate dehydrogenase (LDH)

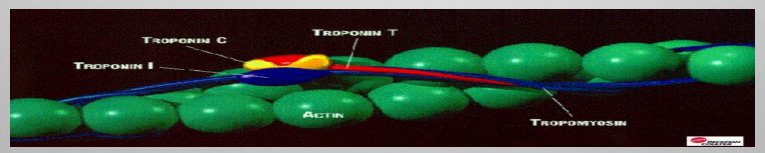
- LDH is a **tetramer**, each chain may be one of two types (H,M) where LDH1 is (H4) while LD5 is (M4)
- LD1 & LD2 predominates in heart
- LDH increases later than CK-MB and Ck
- Reaches a max. level in 48 h
- Remains elevated for 5-6 days after the MI



- A non-specific marker of tissue injury: High levels are found in liver, lung, kidney and other diseases.
- Myocardial infarction resulting in insufficient oxygen delivery to that portion of cardiac muscle. This causes the affected muscle to rely on <u>anaerobic</u> <u>metabolism</u> for its energy supply with concomitant production of lactic acid.

TROPONIN

- TROPONIN IS A **PROTEIN**.
- PRESENT IN HIGH CONCENTRATION IN MUSCLE & HEART.
- REGULATES THE FORCE OF MUSCULAR CONTRACTIONS
- IS COMPOSED OF 3 SUB UNITS I, T AND C.
- TROPONIN C: CA⁺⁺ BINDING. (NOT HEART-SPECIFIC).
- THE **TROPONIN I AND TROPONIN T** FOUND IN <u>HEART</u>
 <u>MUSCLE</u> IS SIGNIFICANTLY **DIFFERENT** FROM TROPONINS
 FOUND IN NON-CARDIAC MUSCLE



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TROPONIN T

- TROPOMYOSIN BINDING ELEMENT.
- ITS LEVEL INCREASES WITHIN 6 HRS OF MI.
- **PEAKS** AT 72 HRS .
- **<u>REMAINS</u>** ELEVATED **7-10 DAYS**.
- TROPONIN T MAY BE ELEVATED IN PATIENTS WITH CHRONIC RENAL FAILURE AND THUS MAY NOT BE SO CARDIAC-SPECIFIC

TROPONIN I:

- IT IS RELEASED WITHIN 4-6 HRS OF THE ONSET OF MI.
- <u>PEAKS</u>14-24HRS.
- **<u>REMAINS</u>** ELEVATED FOR 3-5 DAYS.
- DISAPPEARS FROM BLOOD AFTER ABOUT ONE WEEK. SO, USEFUL FOR DIAGNOSIS OF <u>DELAYED ADMISSION CASES.</u>
 - CARDIAC TROPONINS HAVE BEEN RECOMMENDED AS THE BIOCHEMICAL CARDIAC MARKER OF CHOICE.

CARDIAC TROPONIN: TROPONIN I (CTN I)

- SERUM TROPONINS ARE NOT FOUND IN HEALTHY INDIVIDUALS (UNLIKE CK/MB).
- TROPONINS ARE BOTH <u>MORE SENSITIVE</u> (DIAGNOSE MINOR INFARCTION) AND <u>MORE SPECIFIC</u> THAN CK-MB IN TERMS OF ITS DIAGNOSTIC ABILITY WITH RESPECT TO MYOCARDIAL DAMAGE.
- **PROGNOSTIC MARKER** (RELATION BETWEEN LEVEL IN BLOOD & EXTENT OF CARDIAC DAMAGE). DETERMINATION OF SIZE OF INFARCT.
- DETERMINATION OF **SUCCESS OF REPERFUSION**.
- TWO NEGATIVE TROPONINS <u>6 HOURS APART</u> ARE GOOD (BUT NOT ABSOLUTE) EVIDENCE OF NO RECENT AMI.
- POSITVE TROPONIN IN PATIENTS WITHOUT ECG CHANGES & WITH NORMAL CK-MB LEVELS MAY IDENTIFY PATIENTS AT INCREASED RISK OF CARDIAC EVENTS

HEART-TYPE FATTY ACID-BINDING PROTEIN (H-FABP)

- H-FABP IS A SMALL CYTOSOLIC PROTEIN FOUND IN THE CARDIAC TISSUES.
- IT IS CHIEFLY PRESENT IN THE MYOCARDIUM AND, TO A LESSER EXTENT, IN THE BRAIN, KIDNEY AND SKELETAL MUSCLE.
- RESPONSIBLE FOR THE TRANSPORT OF FATTY ACIDS FROM THE PLASMA MEMBRANE TO:
- SITES OF B-OXIDATION IN **MITOCHONDRIA** AND **PEROXISOMES**.
- > ENDOPLASMIC RETICULUM FOR LIPID SYNTHESIS.
- H-FABP IS RELEASED EXTREMELY EARLY INTO THE SERUM FOLLOWING MYOCYTE RUPTURE.
- >↑↑↑ AS EARLY AS <u>30 MIN</u> AFTER MYOCARDIAL INJURY
- > PEAKS AT <u>6-8 H</u> AND
- **RETURNS** TO BASELINE LEVELS AT <u>~24 H.</u>
- IT COULD BE USED TO QUICKLY **<u>RULE OUT</u>**AMI.

ISOENZYME BB GLYCOGEN PHOSPHORYLASE (GPBB)

- IT IS ONE OF THE 3 ISOFORMS OF GLYCOGEN PHOSPHORYLASE.
- GPBB EXISTS IN CARDIAC AND BRAIN TISSUE. BECAUSE OF THE BLOOD-BRAIN BARRIER, GP-BB CAN BE SEEN AS BEING SPECIFIC TO HEART MUSCLE.
- ↑↑↑ BLOOD LEVELS CAN BE SEEN IN ISCHEMIA, MI AND UNSTABLE ANGINA.
- EARLY BIOCHEMICAL MARKER OF MYOCARDIAL NECROSIS.
- ↑↑↑ WITHIN THE **FIRST HOUR** OF **MI**.
- ↑↑↑ 1–3 HOURS AFTER ISCHEMIA.
- VERY SENSITIVE INDICATOR OF MI WITH A SENSITIVITY SUPERIOR TO THAT OF MYOGLOBIN, CK-MB MASS, AND CTNT.

COPEPTIN

 COPEPTIN, THE C-TERMINAL PORTION OF PROVASOPRESSIN IS COSECRETED WITH VASOPRESSIN.

• **↑↑↑** WITHIN MINUTES IN PATIENTS WITH **AMI**.

• ADDING COPEPTIN + CTNI CAN RULE OUT OF AMI.

ISCHEMIA-MODIFIED ALBUMIN (IMA)

- IT IS RAISED IN THE PRESENCE OF MYOCARDIAL ISCHEMIA.
- NORMAL ALBUMIN CAN BIND METALS AT ITS N TERMINUS.
- DURING ISCHEMIA, FREE RADICALS, ALTER THE BINDING SITE, DECREASING BINDING ABILITY MAKE IT MORE RESISTANT TO BIND METALS.
- **POSITIVE TEST** ISCHEMIA
- NEGATIVE TEST (TOGETHER WITH NEGATIVE TROPONIN AND NEGATIVE ECG) HAS A 99% NEGATIVE PREDICTIVE VALUE FOR MI.
- RAPIDLY CLEARED
- NOT SPECIFIC FOR CARDIAC ISCHEMIA.
- IT IS A MARKER SENSITIVE FOR ISCHEMIA RATHER THAN NECROSIS.
- ➢ IT IS <u>DETECTED</u> WITHIN A FEW MINUTES.
- PEAKS AT 2-4 HOURS.
- ➢ <u>DISAPPEARS</u> WITHIN 6 HOURS.

MICRO-RNAS

MICRO RNA (MIRNA)

- MICRORNAS (MIRNAS) CIRCULATE IN THE BLOODSTREAM IN A REMARKABLY STABLE FORM.
- BECAUSE OF THEIR STABILITY AND OFTEN TISSUE- AND DISEASE-SPECIFIC EXPRESSION AND THE POSSIBILITY TO MEASURE THEM WITH HIGH SENSITIVITY AND SPECIFICITY, MIRNAS ARE EMERGING AS NEW DIAGNOSTIC & PROGNOSTIC BIOMARKERS.

DIAGNOSTIC ABILITIES OF CIRCULATING MIRNAS FOR MI.

- IT HAS BEEN FOUND THAT <u>MIR-1, MIR-133</u>, AND <u>MIR-499</u> WERE
 ELEVATED IN PATIENTS WITH MI.
- THE SLOW TIME COURSE OF <u>MIR-499</u> MIGHT LEAD TO INCREASED DIAGNOSTIC PERFORMANCE AT LATE TIME POINTS AFTER MI WHEN₈
 CTNI HAS ALREADY RETURNED BACK TO NORMAL LEVELS.

MICRO RNA (MIRNA)

DIAGNOSTIC ABILITIES OF CIRCULATING MIRNAS FOR MI.

- _THE **CARDIAC-SPECIFIC** <u>MIR-208</u> WAS NOT DETECTABLE IN PLASMA OF HEALTHY CONTROLS OR IN PATIENTS WITH STABLE CAD.
- WITHIN 4 H AFTER THE ONSET OF SYMPTOMS, MIR-208 WAS DETECTED IN ALL PATIENTS, WHEREAS CTNI WAS ONLY DETECTED IN 85% OF THE PATIENTS, CONFIRMING THE SUPERIOR SENSITIVITY OF MIR-208 AT EARLY TIME POINTS.

DIAGNOSTIC ABILITIES OF CIRCULATING MIRNAS FOR MI.

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 MIR-122 AND MIR-375 EXPERIENCED A <u>DROP</u> IN THEIR PLASMA LEVELS FOLLOWING MI.

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IT MAY BE EXPECTED THAT IN THE FUTURE, A PANEL OF MIRNAS, PROBABLY IN COMBINATION WITH CTNI, HAS A BETTER POTENTIAL TO OFFER SENSITIVE AND SPECIFIC DIAGNOSTIC TESTS FOR AMI.

SALIVARY BIOMARKERS ASSOCIATED WITH MI

- **SALIVA** OFFERS AN EASY, SIMPLE AND NON-INVASIVE PROCEDURE.
- WHOLE SALIVA CONTAINS CONSTITUENTS FROM SERUM, GINGIVAL FLUID AND ORAL MUCOSAL TRANSUDATE.

SALIVARY MARKERS OF ACUTE MYOCARDIAL INFARCTION:

- MYELOPEROXIDASE (MPO), C-REACTIVE PROTEIN (CRP), MYOGLOBIN, CK-MB AND CTN.
- SALIVA CAN BE USED AS AN ALTERNATIVE TO SERUM IN THE DIAGNOSIS OF MI.

RECENTLY: USING NANOCHIPS AND A SWAB OF THE CHEEK,

CARDIAC BIOMARKER READINGS FROM SALIVA WITH ECG READINGS

DETERMINE WITHIN MINUTES WHETHER SOMEONE HAD A HEART ATTACK.



