

HEART FAILURE

Hanna K. AL-Makhamreh, MD FACC

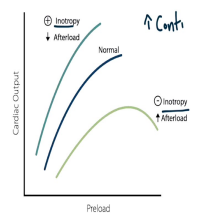
Associate Professor of Cardiology

University Of Jordan

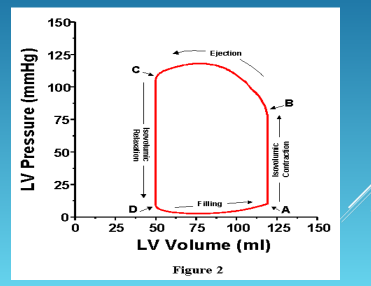
PHYSIOLOGY (FRANK-STARLING) CURVE

- Preload reduction
 - Diuretics
 - venodilators
- Vasodilators ACEI
- Inotropes
 Dobutamine

Starling Curve



PRESSURE-VOLUME LOOP

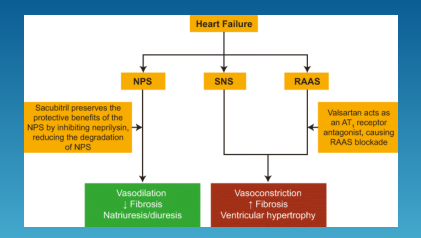


PATHOPHYSIOLOGY

Initial Compensation for impaired myocyte contractility:
Frank-Starling mechanism
Neurohumoral activation

• intravascular volume

•Eventual decompensation •ventricular remodeling •myocyte death/apoptosis •valvular regurgitation

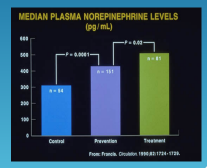


PATHOPHYSIOLOGY OF HF

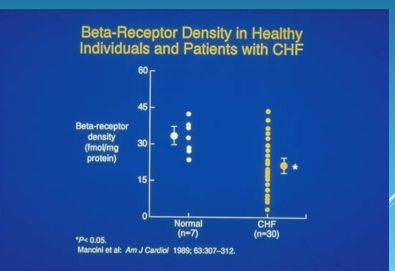
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PATHOPHYSIOLOGY: NEUROHUMORAL

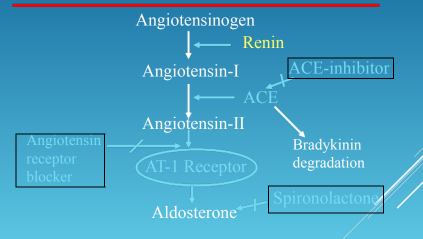
- Adrenergic nervous
 system
- Renin-angiotensinaldosterone system
- Natriuretic peptides



NEUROHUMORAL



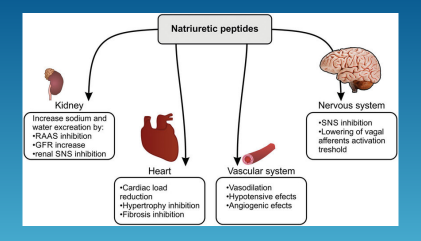
RENIN-ANGIOTENSIN-ALDOSTERONE PATHWAYS



ANGIOTENSIN-II EFFECTS

- Vasoconstriction
- Aldosterone production
- Myocyte hypertrophy
- Fibroblast proliferation
- Collagen deposition

- Apoptosis
- Pro-thrombotic
- Pro-oxidant
- Adrenergic stimulation
- Endothelial dysfunction



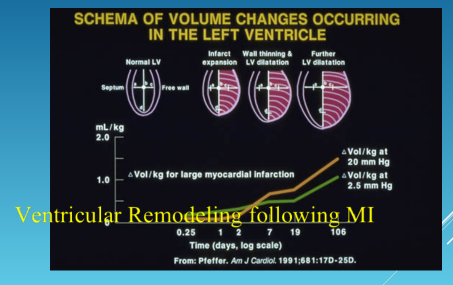
NP

10

THE KIDNEY AND THE HEART FAILURE

- Reduced renal blood flow
- Reduced glomerular filtration rate
- Increased renin production
- Increased tubular sodium reabsorption
- Increased free water retention (vasopressin)

VENTRICULAR REMODELING IN HEART FAILURE



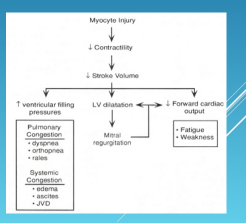
CLINICAL FINDINGS

Biventricular Congestive Heart Failure

-Low forward Cardiac Output -fatigue, lightheadedness, hypotension

-Pulmonary Congestion -Dyspnea, -orthopnea, & PND

-Systemic Congestion -Edema -Ascites -Weight gain



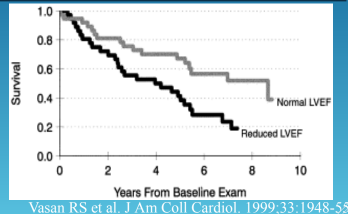
PHYSICAL EXAM

Decreased C.O. Tachycardia BP and pulse pressure Pulsus Alternans (end-stage) Pulmonary venous congestion: laterally displaced PMI

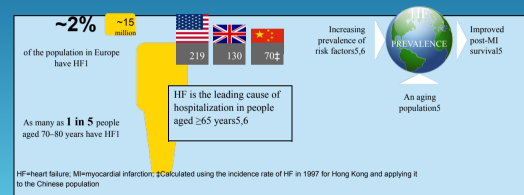
DIAGNOSTIC STUDIES

CXR -enlarged cardiac silhouette, vascular redistribution interstitial edema, pleural effusions EKG –normal tachycardia, atrial and ventricular enlargement, LBBB, RBBB, Q-waves Blood Tests (KFT, BNP, ANA, RF, Fe2+, TFT's, ferritin,) **Echocardiography** LV size, wall thickness function **Cardiac Catheterization**

INFLUENCE OF EF ON SURVIVAL IN PATIENTS WITH HEART FAILURE



HF IS A MAJOR AND GROWING PUBLIC HEALTH PROBLEM



1. Dickstein et al. Eur Heart J 2008;29:2388–442; 2. Go et al. Circulation 2013;127:e6–e245; 3. Allender et al. Coronary Heart Disease Statistics 2008; 4. Hung et al. Hong Kong Med J 2000;6:159–62; 5. Hunt et al. J Am Coll Cardiol 2009;53:e1–90; 6. Kearney et al. Lancet 2005; 365:217–23; 5. Forman et al. Am Heart J 2009;175:1010–17; 6. Healthcare Cost and Utilization Project 2009 (http://www.hcun-us.ahur.org/project/factsandfoures/2009/CDC_ 2009 isp.Accessed_lanuary 2013

HF IMPOSES A SIGNIFICANT ECONOMIC BURDEN ON THE HEALTHCARE SYSTEM



\$\$\$\$\$\$\$\$ \$\$\$\$\$\$\$ \$\$\$\$\$\$\$ 10 %

OF THE COST OF HF IS DUE TO HOSPITALIZATIONS

OF THE COST OF HF IS DUE TO PHARMACOLOGICAL TREATMENT2

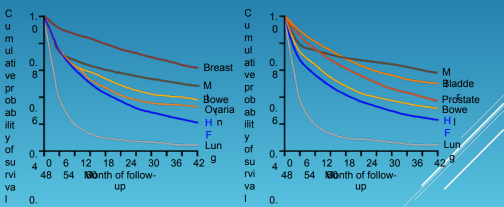
HF=heart failure; ‡USA estimate includes direct costs (total annual medical spending) and indirect costs (lost productivity due to morbidity and mortality)

1. Dickstein et al. Eur Heart J 2008;29:2388–442; 2. Hunt et al. J Am Coll Cardiol 2009;53:e1–90; 3.Go et al. Circulation 2013;127:e6–e245

MORTALITY FOLLOWING ADMISSION FOR ACUTE HEART FAILURE EXCEEDS THAT OF MOST CANCERS Female survival rates (%): Male survival rates (%):

HF, MI and other malignancies

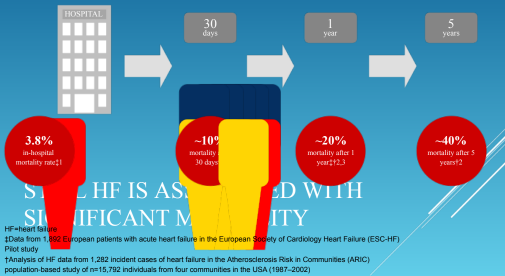
Male survival rates (%): HF, MI and other malignancies



2All patients with a first admission to any Scottish hospital in 1291 for HF, MI or the four most common types of cancer specific to men and women were identified, and 5-year survival rates compared

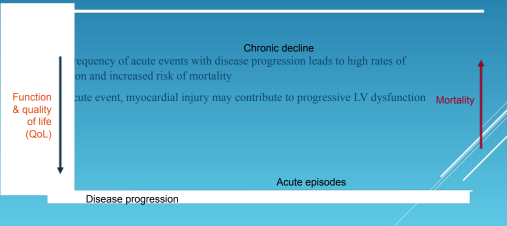
Stewart et al. Eur J Heart Fail 2001;3:315-

0



1. Maggioni et al. Eur J Heart Fail 2010;12:1076–84; 2. Loehr et al. Am J Cardiol 2008;101:1016–22; 3. Maggioni et al. Eur J Heart Fail 2013;15:808–17

HEART FAILURE IS A PROGRESSIVE CONDITION WITH HIGH MORBIDITY AND MORTALITY



Gheorghiade et al. Am J Cardiol 2005;96:11G–17G; Gheorghiade & Pang. J Am Coll Cardiol 2009;53:557–73



- Most patients with HF experience symptoms due to impaired LV myocardial function1
- The most common causes of HF are coronary heart disease (CHD), valve disease and cardiomyopathies2

*Including hypertension, diabetes, exposure to cardiotoxic agents, peripartum cardiomyopathy, etc.

CHD is the underlying cause of 60–70% of acute HF cases3

1. Hunt et al. J Am Coll Cardiol 2009;53:e1-90

2. Dickstein et al. Eur Heart J 2008;29:2388-442

3. Nieminen et al. Eur Heart J 2005;26:384-416

HIGH PREVALENCE OF MULTIPLE CO-MORBIDITIES

Many patients with chronic HF have a range of co-morbidities that contribute to the cause of the disease and play a key role in its progression and in the response to therapy

hypertension*

ischemic heart disease*

diabetes mellitus

cardiac arrhythmias

ventricular arrhythmias

atrial fibrillation

respiratory disorders

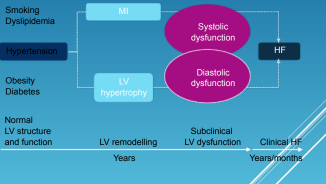
cognitive dysfunctio

hyperlipidemia

chronic anemia

renal failure

arthritis



*Major contributors to development of HF

This can lesul in patients buildened with multiple pills per day, each with different dosage

GUIDELINE DEVELOPMENT

ACCF-AHA 2013

ESC 2012

HFSA 2010

NICE AHF 2014/ CHF 2010

| Level of Evidence | | Class of Recommendation | | |
|-------------------|--|-------------------------|---|--|
| | Multiple populations evaluated* Data from multiple randomized | I | Benefit >>> Risk Procedure/Treatment SHOULD be performed/administered | |
| | clinical trials or meta-analyses | Ha | Benefit >> Risk (Additional studies with focused objectives needed) IT IS REASONABLE to perform procedure/administer treatment | |
| | Data from single randomized clinical trial or nonrandomized studies | IIb | Benefit ≥ Risk (Additional studies with broad objectives needed; additional registry data would be helpful) Procedure/Treatment <i>MAY BE CONSIDERED</i> | |
| | Very limited populations evaluated* Consensus of opinion of the experts, case studies, or standard-of-care | ш | No Benefit: Procedure/test is not helpful and treatment has no proven benefit Harm: Procedure/test is expensive without benefit or harmful and treatment is potentially harmful to patients | |

HEART FAILURE DEFINITION

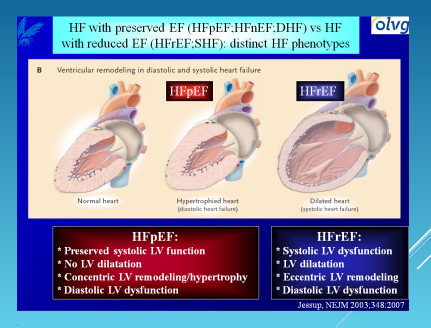
| Heart Failure | |
|---------------|--|
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| | |
| | |

AHF, acute heart failure; HF, heart failure

| Progression IIFrEF ≤40% ≤35% <50% | ased on Clinical | Types | ACCF-AHA 2013 | ESC 2012 | HFSA 2010 | NICE 201 |
|--|------------------|-------|--------------------------------------|---|-----------|-----------------------------------|
| Status • 41%-49% (mrHF,) • 35-50% 'grey area'; most No three of LN of | | HFrEF | <u>≤40%</u> | ≤35% | <50% | |
| improved(HFiEF) systolic dysfunction | | | 41%-49% (mrHF,) ≥10% to be ≥40%, | 35–50% 'grey area'; most probably have primarily mild | ≥50% | No threshol of LVEF defined |

HFrEF, heart failure with reduced ejection fraction; HFpEF, heart failure with preserved ejection fraction; LVEF, left ventricular ejection fraction

HEART FAILURE SUBTYPES



| Based on the LVEF | | |
|----------------------------------|-------|--|
| Based on the Functiona Status | | |
| | Class | Severity of symptoms and limitation of physical activity |
| Based on Clinical Progression | 1 | No limitation of physical activity |
| Rased on Hemodynam | | Ordinary physical activity does not cause symptoms of HF (breathlessness, fatigue, or palpitations) |
| Status | п | Slight limitation of physical activity |
| | | Comfortable at rest, but ordinary physical activity results in symptoms of HF |
| | Ш | Marked limitation of physical activity |
| | | Comfortable at rest, but less than ordinary physical activity causes symptoms of HF* |
| | IV | Unable to carry on any physical activity without discomfort/symptoms of HF, or symptoms of HF at rest may be present |
| | | If any physical activity is undertaken, discomfort is increased |
| | | further classify class III into IIIA (comfortable at rest, but less than ordinary physical activity causes symptoms of HF) and IIIB st, but minimal exertion causes fatigue, palpitation, or dyspnea) |

| Based on the LVEF | | | |
|------------------------------------|--------------|---|-----------------------------|
| Based on the Functiona Status | | | |
| Based on Clinical | | | |
| Progression Based on Hemodynami | Stages of HF | Development and progression of HF | Corresponding NYHA Class |
| Status | А | At high risk for HF but without structural heart disease or symptoms of HF $$ | None |
| | В | Structural heart disease but without signs or symptoms of HF | I |
| | | | I |
| | с | Structural heart disease with prior or current symptoms of HF | П |
| | | | ш |
| | D | Refractory HF requiring specialized interventions | IV |
| | | | |

HF, heart failure; NYHA, New York Heart Association

| Based on the LVEF Based on the Function Status Based on Clinical | | | | |
|---|--|----|--|--------------------------|
| Frenzression Based on Hemodynami Status | | | Congestic (e.g. orthopnea, elevated pulmonary rales, | jugular venous pressure, |
| | | | No | Yes |
| | Low perfusion at rest? (e.g. narrow pulse pressure, cool extremities, hypotension) | No | Warm and Dry | Warm and Wet |
| | carcinetes, nypotension) | | Cold and Dry | Cold and Wet |
| | | | | |

HF, heart failure

SYMPTOMS



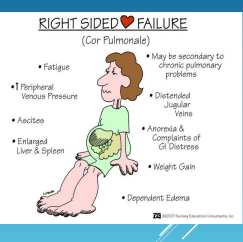










Figure 24. CXR Showing Acute Decompensated Heart Failure





| Method | ESC* | Purpose |
|----------------|------|---|
| ECG | IC | Shows the heart rhythm and electrical conduction. Important for decisions about treatment (e.g. rate control and anticoagulation for AF, pacing for bradycardia, or CRT if the patient has LBBB). It may show evidence of LV hypertrophy or Q waves (indicating loss of viable myocardium), giving a possible clue to the etiology of HF. |
| Chest X-ray | | Most useful in identifying an alternative, pulmonary explanation for a patient's symptoms and signs. It may show pulmonary venous congestion or edema in a patient with HF. |
| Echocardiogram | IC | Provides immediate information on chamber volumes, ventricular systolic and diastolic function, wall thickness, and valve function. |

The echocardiogram and electrocardiogram are the most useful tests in patients with suspected HF

McMurray et al. Eur Heart J 2012:33:1787–847

INVESTIGATIONS TO CONSIDER IN SELECTED PATIENTS LABORATORY TESTS

| Method | ESC* | Purpose |
|---|------|---|
| Biochemical and hematological investigations | IC | Determine whether RAAS blockade can be initiated safely (renal function and potassium). Exclude anemia (can mimic or aggravate HF). |
| Natriuretie Peptide (NP) | | Where the availability of echocardiography is limited, an alternative approach to diagnosis is to measure the blood concentration of NP. NP levels also increase with age, renal insufficiency, but may be reduced in obese patients. A normal NP level in an untreated patient virtually excludes significant cardiac disease, making an echocardiogram unnecessary. |

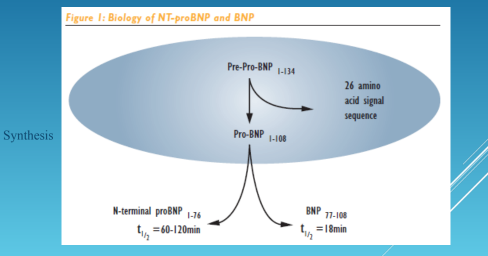
*ESC recommendation, class and level of evidence

NP: natriuretic peptide; RAAS: renin-angiotensin-aldosterone system

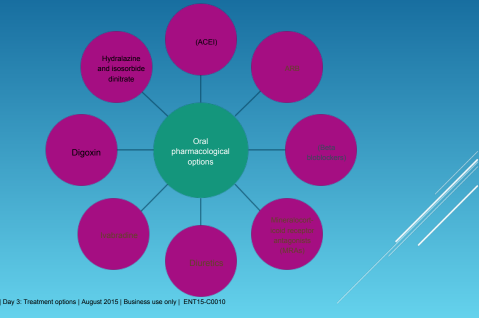
CARDIAC NATRIURETIC PEPTIDES

- What is BNP?
 - A 32 amino acid polypeptide
 - Belong to a class of structurally similar natriuretic peptides (classes A,B,C and D)
 - Secreted by cardiac myocytes (mainly left) in response to excessive distension of the Heart ventricles
 - Similar to ANP (Atrial Natriuretic Peptide) but has longer t1/2 (~20mins, double that of ANP) Named after extracts found in Pig-brain
- What is NT-proBNP?
 - NT-proBNP is a biologically inactive 76 amino acid N-terminal fragment
 - · Co-secreted with BNP
 - Even longer t1/2 than BNP (~1-2hrs vs ~20mins)
- Biological effects of Cardiac Natriuretic peptides
 - Increase Natriuresis

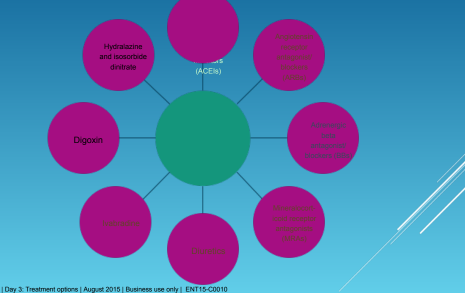
SYNTHESIS IN MYOCYTES



WHAT ARE THE ORAL PHARMACOLOGICAL OPTIONS?

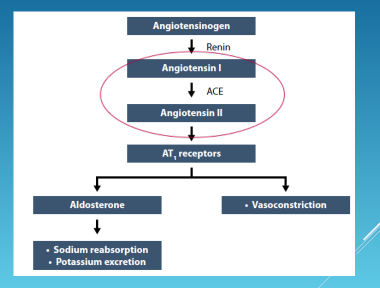


WHAT ARE THE ORAL PHARMACOLOGICAL OPTIONS?



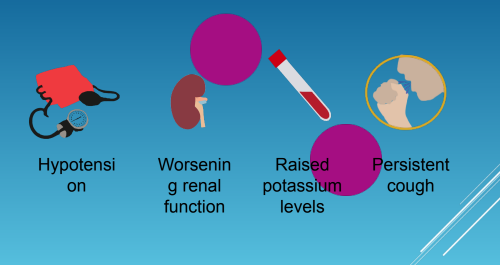
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ACEIS: HOW THEY WORK - RAAS

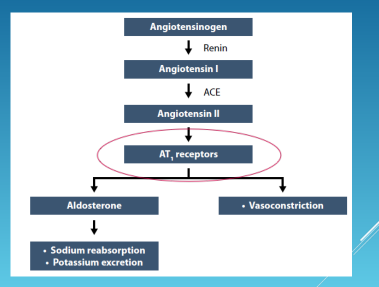


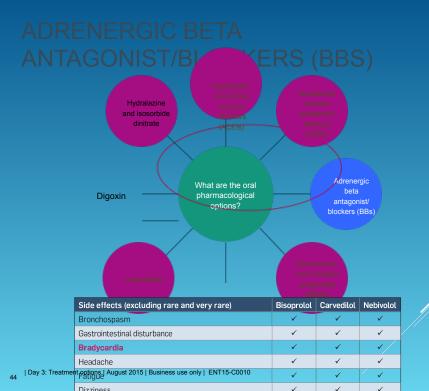
| Day 3: Treatment options | August 2015 | Business use only | ENT15-C0010

ACEIS: RISKS



ARBS: HOW THEY WORK - RAAS

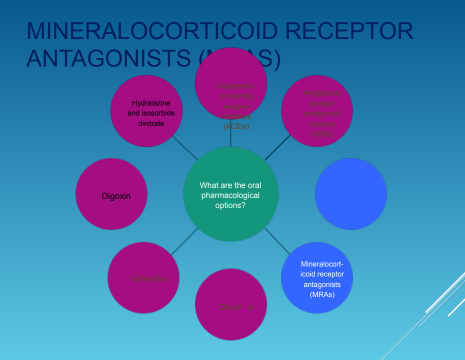




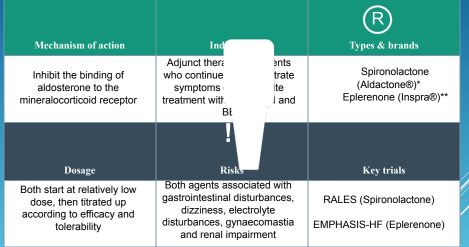
BETA BLOCKERS: RISKS (1)

| Side effects (excluding rare and very rare) | Bisoprolol | Carvedilol | Nebivolol |
|--|------------|------------|-----------|
| Bronchosp | ✓ | ✓ | ✓ |
| Gastroin | ✓ | ~ | ✓ |
| | 1 | ✓ | ✓ |
| Headach | ✓ | ~ | ✓ |
| Fatigue | ✓ | ~ | ~ |
| Dizziness | 1 | ~ | ~ |
| Paraesthesia | ✓ | ~ | ✓ |
| Heart failure | ✓ | | ✓ |
| Hypotension | 1 | ~ | ✓ |
| Conduction disorders | ✓ | | ✓ |
| Peripheral vasoconstriction, e.g. claudication and Raynaud's | ~ | | ~ |
| Dyspnoea | ✓ | ð | ~ |
| Sleep disturbances | ✓ | | ~ |
| Vertigo | 1 | | ~ |
| Psychosis | 1 | | ~ |
| Sexual dysfunction | ✓ | | 1 |
| | | | |

δ Postural hypotension. Δ Exacerbation of previous condition. Π Also eye irritation. J Also painful extremities.



MINERALOCORTICOID ANTAGONISTS (MRAS): THE FACTS

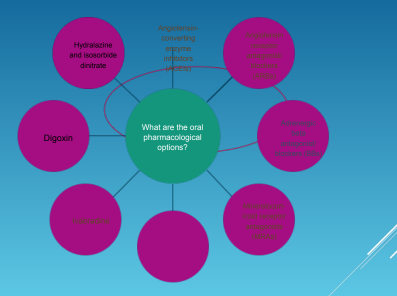


*A non-proprietary drug is available

** A non-proprietary drug is not available

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DIURETICS



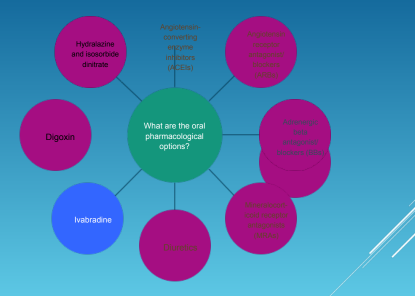
DIURETICS: THE FACTS

| | | | R |
|--|--|----------------------|--|
| Mechanism of action | Indi | 1. | Bendroflumenthiazide |
| Thiazide diuretics - inhibit the reabsorption of sodium in the kidney's distal convoluted tubule Loop diuretics - inhibit absorption from the kidney's loop of Henle | Patients wit deemed to hav、 | are ,erload 4. | (thiazide) (Aprinox®, Neo- Naclex®)* Chlortalidone (thiazide- related) (Hygroton®)** Furosemide (loop) (Rusyde®, Frusol®)* Bendroflumenthiazide (loop)(Torem®)* |
| Dosage | Risks | | Key trials |
| Bendroflumenthiazide: 5-10 mg daily Chlortalidone: 25-30 mg daily Furosemide: 40 mg mg daily Bendroflumenthiazide: 5 mg daily | Both types of diuretics associated with mild gastrointestinal side effects, postural hypotension, metabolic and electrolyte disturbances, blood disorders | | Paucity of trial evidence for the efficacy of diuretics in HF. They are recommended for their beneficial effects on dyspnoea and oedema |

*A non-proprietary drug is available ** A non-proprietary drug is not available

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IVABRADINE



• Acts as a specific bradycardic agent, lowers heart rate by specific action on the sino-atrial node controlled by If current without affecting other cardiac ionic currents. It has no negative inotropic effect and has beneficial effects on left-ventricular systolic dysfunction. The only negative effects are vision disturbances which are mild and transient.

• Ivabradine is the first selective sinus node If channel inhibitor that results in a decrease in the slope of the diastolic depolarization in the SA node cells

- It is rapidly and almost completely absorbed after oral administration with a peak plasma level reached in approximately 1 hour under fasting condition.
- The absolute bioavailability of the 10mg dose is around 40%
- No side effects like sexual disturbances, respiratory side effects, bradycardia or rebound phenomena

- Indication
- Angina pectoris (2005) CHF (2012 in EU, 2015 in US); for use in heart failure patients inadequately controlled with optimal dose of beta-blocker (or intolerant) and whose heart rate is >75 bpm in EU and ≥70 bpm in US

DIGOXIN



DIGOXIN

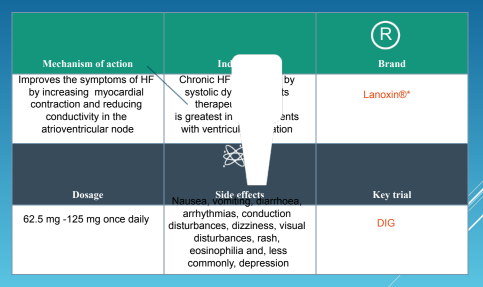
Cardiac glycoside

Addresses heart failure symptoms by increasing myocardial contraction and reducing conductivity in atrioventricular node

Generally considered for patients with persistent symptoms

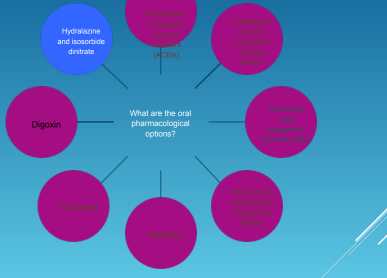
Despite other treatments - ACEI and BB + other agents e.g. spironolactone, ARB, or hydralazine/nitrate

DIGOXIN: THE FACTS

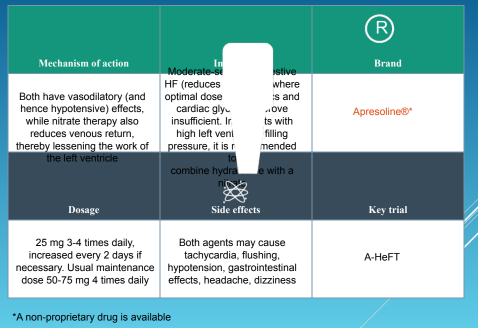


*A non-proprietary drug is available

HYDRALAZINE AND ISOSORBIDE



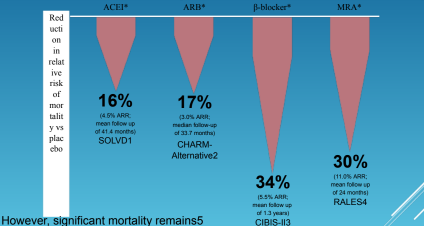
HYDRALAZINE AND ISOSORBIDE DINITRATE: THE



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SUCCESSFUL INTERVENTION BY ADRESSING NEUROHORMONAL ACTIVIATION

Chronic HFrEF survival rates have improved over time with the introduction of new therapies

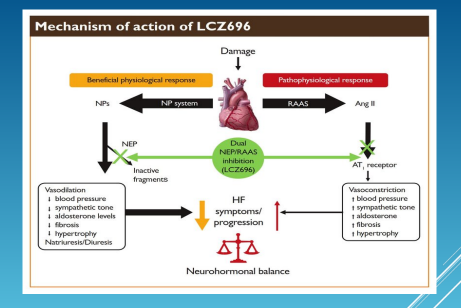


*On top of standard therapy at the time of study (except in CHARM-Alternative where background ACEI therapy was excluded). Patient populations varied between trials and as such relative risk reductions cannot be directly compared. SOLVD (Studies of Left Ventricular Dysfunction), CBIS-II (Cardiac Insufficiency Bisoprolol Study II) and RALES (Randomized Aldactone Evaluation Study) enrolled chronic HF patients with LVEFS35%. CHARM-Alternative (Candesartan in Heart failure: Assessment of Reduction in Mortality and Morbidity) enrolled chronic HF patients with LVEFS40%.

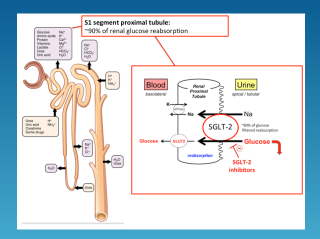
ARR=absolute risk reduction; MRA=mineralocorticoid receptor antagonist; RRR=relative risk reduction

1. SOLVD Investigators. N Engl J Med 1991;325:293-302; 2. Granger et al. Lancet 2003;362:772-6

3. CIBIS-II Investigators. Lancet 1999;353:9–13; 4. Pitt et al. N Engl J Med 1999;341:709-17; 5. Roger et al. JAMA 2004;292:344–50



ARNI (SACUBITRIL/VALSARTAN)



SGLT2 INHIBITORS

CHF – LEVEL OF RECOMMENDATIONS

Re

| Drug Classes | Pharmacological therapies | ACCF- AHA 2013 | HFSA 2010 | ESC 2012 | NICE CHF-2010 |
|---------------------------|---|-----------------------------|--|---------------------------------|-----------------------------------|
| Level of commendations | ARNI/ACEI/ARB | IA | А | IA | А |
| (1/2) | Beta blockers | IA | А | IA | А |
| CCF-AHA 2013 | Loop diuretics | IC | А | - | С |
| | ARBs | | | | |
| ESC 2012 | In patients who are intolerant to ACEI | IA* | А | IA | Α |
| | In patients with persisting symptoms despite treatment with ACEI and BB, who are intolerant MRA | Hb A | - | IA | - |
| HFSA 2010 | Patients with persisting symptoms despite treatment with ACEI and a beta-blocker | - | А | - | † |
| NICE 2010 | Individual ARBs may be considered as initial therapy rather than ACEI for HF patients post-MI | - | А | - | - |
| | MRAs | | | | |
| | - Patients with persisting symptoms and EF ${\leq}35\%$, despite treatment with an ACEI and beta-blocker | - | A‡ | IA | A# |
| | TARBs are used as alternatives to ACEI (level of recommendation IIa A), tNo of evidence A), for patients with moderate HF (strength of evidence B); #esp | t graded; ‡ in patie | nts with severe HF with moderate to s | or post-MI HF evere HF, NYH/ | (strength A class III- |
| | IV or has had an MI within the past month; **MRAs are recommended in patien | ts with NYHA clas | s III-IV, LVEF<35% | | |

ACEI, angiotensin converting enzyme inhibitor, ARB; angiotensin receptor blocker; BB, beta blockers; EF, ejection fraction; HFrEF, heart failure and reduced ejection fraction; HF, heart failure; MI, myocardial infarction; MRA, mineralocorticoid receptor antagonist; NYHA, New York Heart Association.

CHF – LEVEL OF RECOMMENDATIONS

| Level of |
|-----------------|
| Recommendations |
| (2/2) |
| |

Drug Classes

ACCF-AHA 2013

ESC 2012

HFSA 2010

NICE 2010

| Pharmacological therapies | ACCF- AHA 2013 | HFSA 2010 | ESC 2012 | NICE CHF-2010 |
|--|----------------------|--------------|-------------|------------------|
| Digoxin | | | | |
| In patients with persisting symptoms despite treatment with ACEI/ARB, BB and MRA | Ha B | B/C* | IIb B | А |
| • In patients with sinus rhythm, EF≤45% who are unable to tolerate a beta-blocker (should be given with ACEI+MRA) | - | - | IIb B | - |
| H-ISDN | | | | |
| In symptomatic African-American patients, NYHA class III-IV, despite optimized standard therapy | IA | A/B† | - | : |
| In patients unable to tolerate an ACEI/ARB due to hyperkalemia or renal dysfunction | Ha B | С | IIb B | А |
| Patients with persisting symptoms despite optimized standard therapy (ACEI/ARB, beta-blocker and MRA) | - | С | IIb B | - |
| Ivabradine | | | | |
| In patients with sinus rhythm with an EF ≤35%, HR ≥70 bpm, and persisting symptoms despite treatment with beta- blocker, ACEI and an MRA | - | - | Ha B | \$# |
| • Patients with sinus rhythm with an EF ≤35% and a HR ≥70 bpm who are unable to tolerate beta-blocker | - | - | Пр С | \$# |

ACEI, angiotensin converting enzyme inhibitor; ARB; angiotensin receptor blocker; BB, beta blockers; EF, ejection fraction; HF, heart failure; HR, heart rate; MRA, mineralocorticoid receptor antagonist; MI, myocardial infarction; NYHA, New York Heart Association;

2021 ESC HF GUIDELINES RECOMMENDATIONS FOR THE MANAGEMENT OF PATIENTS WITH HFREF

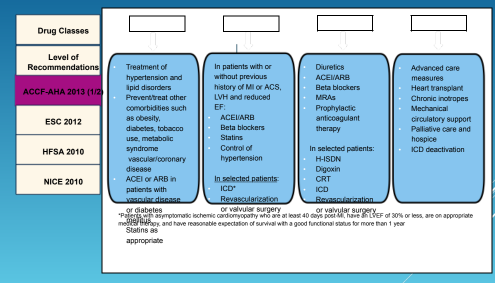
Management of patients with HFrEF1

The 2021 ESC HF Guidelines now recomme

Pharmacological treatments indicated in patients with HFrEF (LVEF ≤40%; NYHA class II–IV)

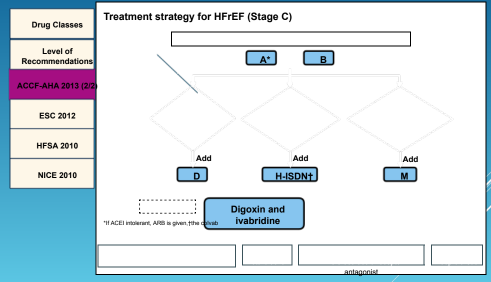
| An ACEi is recommended for patients with HFrEF to reduce the risk of HF hospitalization and death | Α | |
|---|---|---|
| A BB is recommended for patients with stable HFrEF to reduce the risk of HF hospitalization and death | | |
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PHARMACOLOGICAL THERAPY – CHF



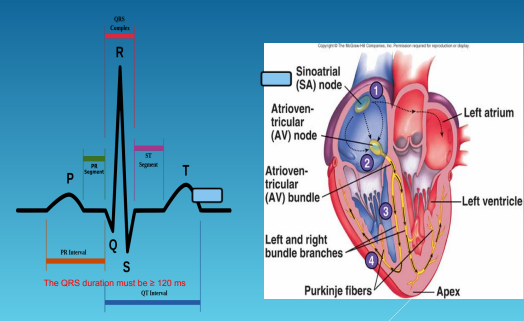
ACEI, anglotensin conventing enzyme inhibitor, ACS, acute coronary syndrome; ARB; anglotensin receptor blocker; CRT; cardiac resynchronization therapy; EF, ejection fraction; H-ISDN, hydralazine and isosorbide dinitrate; ICD, implanatable cardioverter-defbrillator; LVH, left ventricular hypertrophy, MI, myocardial infarction; MRA, mineralocorticid receptor antagonist

PHARMACOLOGICAL THERAPY – CHF



ACEI, angiotensin converting enzyme inhibitor; ARB, angiotensin receptor blocker; Cr., creatinine; HFrEF, heart failure and reduced ejection fraction; NYHA, New York Heart Association

LBBB



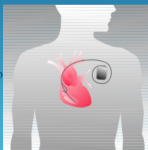
Device implantable inside the body, able to perform both cardioversion, defibrillation and pacing of the heart Indications

- Ventricular tachycardia and ventricular fibrillation (Secondary prevention)
- Prevention of sudden cardiac death (SCD).Patients with an EF <35% (Primary prevention)

CRT: CARDIAC RESYNCHRONIZATION THERAPY

1. Improved hemodynamics

- Increased CO
- Reduced LV filling pressures
- Reduced sympathetic activity
- Increased systolic function w/o MVO2
- 2. Reverse LV remodeling/architecture
 - Decreased LVES/ED volumes
 - Increased LVEF



CRT INDICATION NYC II-IV WITH LBBB

- Improved exercise tolerance
- Reduce symptoms
- Reduced remodeling
- Reduced mortality
- Reduce need for hospitalization rhythm

THANK YOU

