

“Problemas emergentes de Saúde Pública:

A propósito da estratégia 90%-90%-90% da OMS (e da realidade do CHS)”

JOSÉ M. D. POÇAS

DIRETOR DO SDI DO CHS HSB SETÚBAL

O SDI 2ª PARTE : NÓS ... E OS OUTROS!!!



O SDI do CHS HSB EPE de Setúbal

José M. D. Poças

Diretor do SDI



POTENCIAIS CONFLITOS DE INTERESSE...

× “Advisory Boards”

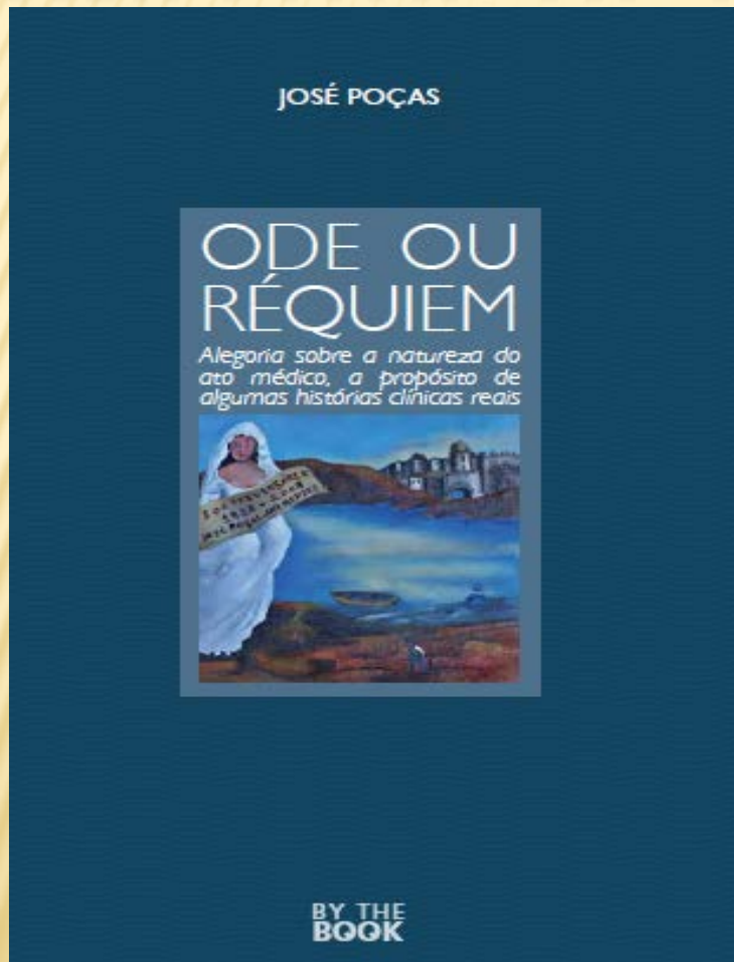
- + Gilead
- + Janssen
- + VIIV
- + BMS
- + MSD



× Congressos e Palestras

- + Abbott
- + Bayer
- + BMS
- + Boheringer Ingelheim
- + Gilead
- + Janssen
- + MSD
- + ROCHE
- + VIIV

OS MEUS ÚNICOS E VERDADEIROS CONFLITOS DE INTERESSE!!!



SUMÁRIO

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- × II)- Os Problemas Emergentes
 - + Multirresistência aos ABs
- × III)- Doenças Emergentes
 - + Infecções transmitidas por vetores e zoonoses
- × IV)- O HIV/SIDA
 - + Os desafios da NOVA estratégia 90/90/90% da ONUSIDA
- × V)- A realidade do CHS
- × VI)- Conclusões

I)- INTRODUÇÃO

O PENSAMENTO DE QUEM SABIA BEM DO QUE FALAVA!!!

- ✘ “... meus senhores, mas são os micróbios que terão a última palavra...” (Louis Pasteur, 1882-1895)

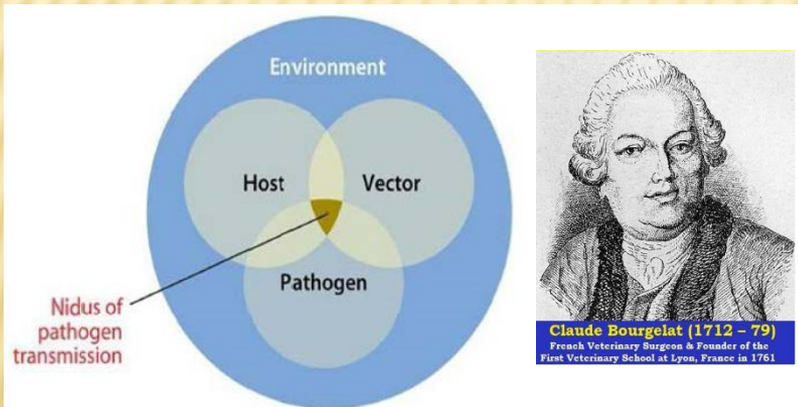
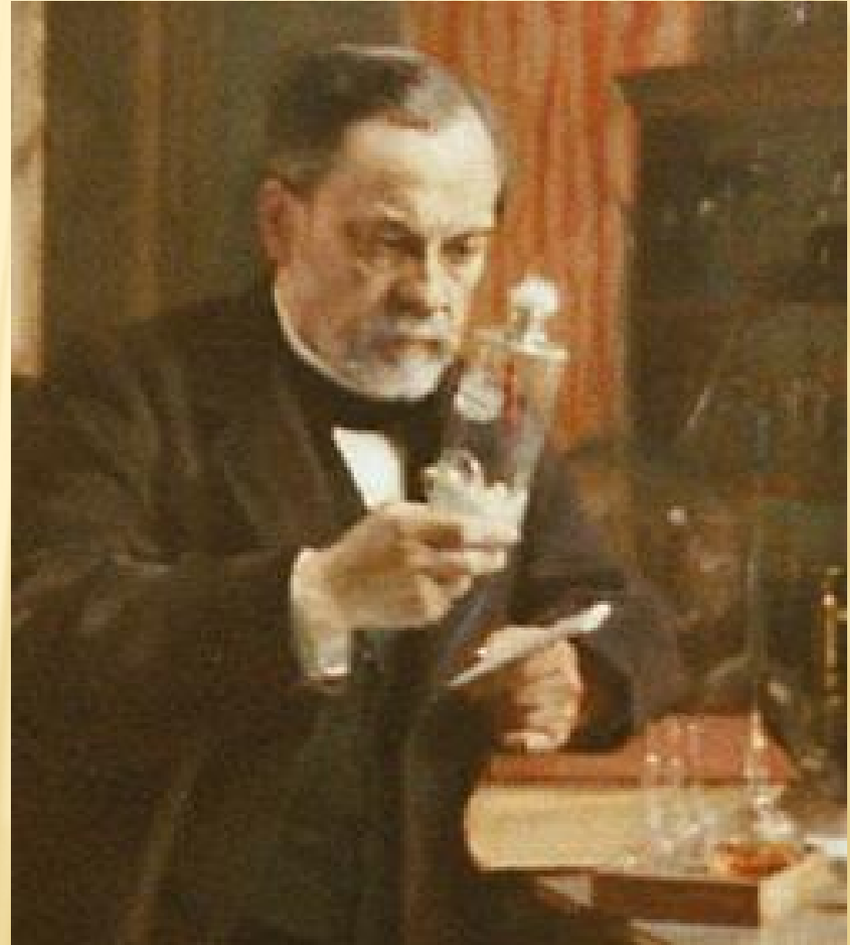


FIGURE WO-3 Key influences on vector-borne plant diseases.
SOURCE: Reisen, 2010. Reproduced with permission of *Annual Review of Entomology*, Volume 55, © by Annual Reviews, <http://www.annualreviews.org>.

APENAS, OS CASOS MAIS RECENTES ...



AS PALAVRAS NÃO PODEM SER UMA MERA CONVENIÊNCIA DE CIRCUNSTÂNCIA

✘ Adenda de 1989

+ “Todo e cada um dos doentes deve beneficiar do melhor tratamento conhecido possível”



O GOVERNO PORTUGUÊS TAMBÉM LEGISLOU NO MESMO SENTIDO

ASSEMBLEIA DA REPÚBLICA

Lei n.º 15/2014
de 21 de março

Lei consolidando a legislação em matéria de direitos e deveres do utente dos serviços de saúde

CAPÍTULO II

Direitos do utente dos serviços de saúde

Artigo 2.º

Direito de escolha

1 — O utente dos serviços de saúde tem direito de escolha dos serviços e prestadores de cuidados de saúde, na medida dos recursos existentes.

2 — O direito à proteção da saúde é exercido tomando em consideração as regras de organização dos serviços de saúde.

Artigo 4.º

Adequação da prestação dos cuidados de saúde

1 — O utente dos serviços de saúde tem direito a receber, com prontidão ou num período de tempo considerado clinicamente aceitável, consoante os casos, os cuidados de saúde de que necessita.

2 — O utente dos serviços de saúde tem direito à prestação dos cuidados de saúde mais adequados e tecnicamente mais corretos.

3 — Os cuidados de saúde devem ser prestados humanamente e com respeito pelo utente.

Artigo 9.º

Queixas e reclamações

1 — O utente dos serviços de saúde tem direito a reclamar e apresentar queixa nos estabelecimentos de saúde, nos termos da lei, bem como a receber indemnização por prejuízos sofridos.

2 — As reclamações e queixas podem ser apresentadas em livro de reclamações ou de modo avulso, sendo obrigatória a resposta, nos termos da lei.

3 — Os serviços de saúde, os fornecedores de bens ou de serviços de saúde e os operadores de saúde são obrigados a possuir livro de reclamações, que pode ser preenchido por quem o solicitar.

CAPÍTULO V

Da Carta dos Direitos de Acesso aos Cuidados de Saúde pelos Utentes do Serviço Nacional de Saúde

Artigo 25.º

Objetivo e conteúdo

1 — A Carta dos Direitos de Acesso visa garantir a prestação dos cuidados de saúde pelo SNS e pelas entidades

convencionadas em tempo considerado clinicamente aceitável para a condição de saúde de cada utente do SNS, nos termos da presente lei.

2 — A Carta dos Direitos de Acesso define:

a) Os tempos máximos de resposta garantidos;

b) O direito do utente à informação sobre esses tempos.

3 — A Carta dos Direitos de Acesso é publicada anualmente em anexo à portaria que fixa os tempos máximos garantidos.

4 — A Carta dos Direitos de Acesso é divulgada no portal da saúde e obrigatoriamente afixada em locais de fácil acesso e visibilidade em todos os estabelecimentos do SNS, bem como em todos os que tenham convencionado a prestação de cuidados de saúde aos seus utentes.

Artigo 26.º

Tempos máximos de resposta garantidos

1 — Para efeitos do disposto no artigo anterior, o membro do Governo responsável pela área da saúde estabelece, por portaria, os tempos máximos de resposta garantidos para todo o tipo de prestações sem carácter de urgência, nomeadamente ambulatório dos centros de saúde, cuidados domiciliários, consultas externas hospitalares, meios **Diário da República, 1.ª série — N.º 57 — 21 de março de 2014 2131** complementares de diagnóstico e terapêutica e cirurgia programada.

2 — Gradualmente, os tempos máximos de resposta garantidos por tipo de prestação são discriminados por patologia ou grupos de patologias.

3 — Cada estabelecimento do SNS, tomando como referência a portaria referida no n.º 1, fixa anualmente, dentro dos limites máximos estabelecidos a nível nacional, os seus tempos de resposta garantidos por tipo de prestação e por patologia ou grupo de patologias, os quais devem constar dos respetivos plano de atividades e contratos -programa.

Artigo 27.º

Informação ao utente

De forma a garantir o direito do utente à informação, previsto no artigo 25.º da presente lei, os estabelecimentos do SNS e do sector convencionado são obrigados a:

a) Afixar em locais de fácil acesso e consulta pelo utente a informação atualizada relativa aos tempos máximos de resposta garantidos por patologia ou grupos de patologias, para os diversos tipos de prestações;

b) Informar o utente no ato de marcação, mediante registo ou impresso próprio, sobre o tempo máximo de resposta garantido para prestação dos cuidados de que necessita;

c) Informar o utente, sempre que for necessário acionar o mecanismo de referenciação entre os estabelecimentos do SNS, sobre o tempo máximo de resposta garantido para lhe serem prestados os respetivos cuidados no estabelecimento de referência, nos termos previstos na alínea anterior;

d) Informar o utente, sempre que a capacidade de resposta dos estabelecimentos do SNS estiver esgotada e for necessário proceder à referenciação para os estabelecimentos de saúde do sector privado, nos termos previstos na alínea b);

e) Manter disponível no seu sítio da Internet informação atualizada sobre os tempos máximos de resposta garantidos nas diversas modalidades de prestação de cuidados;

f) Publicar e divulgar, até 31 de março de cada ano, um relatório circunstanciado sobre o acesso aos cuidados que prestam, os quais serão auditados, alentoria e anualmente, pela Inspeção -Geral das Atividades em Saúde.

Artigo 28.º

Reclamação

É reconhecido ao utente o direito de reclamar para a Entidade Reguladora da Saúde (ERS), nos termos legais aplicáveis, caso os tempos máximos garantidos não sejam cumpridos.

Aprovada em 20 de fevereiro de 2014.

A Presidente da Assembleia da República, **Maria da Assunção A. Esteves**.

Promulgada em 11 de março de 2014.

Publique -se.

O Presidente da República, **ANIBAL CAVACO SILVA**.

Referendada em 13 de março de 2014.

O Primeiro -Ministro, **Pedro Passos Coelho**.

O MAIS ECONÓMICO DOS MELHORES TRATAMENTOS E NÃO O MELHOR DOS MEDICAMENTOS MAIS BARATOS



CONSELHO NACIONAL DE ÉTICA PARA AS CIÊNCIAS DA VIDA

64/CNECV/2012

CONSELHO NACIONAL DE ÉTICA
PARA AS CIÊNCIAS DA VIDA

**PARECER SOBRE UM MODELO DE DELIBERAÇÃO
PARA FINANCIAMENTO DO CUSTO
DOS MEDICAMENTOS**



CNECV

CONSELHO NACIONAL DE ÉTICA PARA AS CIÊNCIAS DA VIDA

D. Conclusões

2. O CNECV recomenda que, nas decisões sobre racionalização de custos, esteja patente que as opções fundamentais serão entre os “mais baratos dos melhores” (fármacos de comprovada efectividade) e não sobre os “melhores dos mais baratos”.

12. Em qualquer caso, o CNECV tem como essencial que tudo o que se faça não pode de modo algum pôr em causa a relação de confiança e de aliança terapêutica entre os doentes e os profissionais de saúde.

II)- OS PROBLEMAS EMERGENTES: **A MULTIRRESISTÊNCIA AOS ABS**

O PRIMEIRO ALERTA!!!

ANNALS OF THE NEW YORK ACADEMY OF SCIENCES
Issue: Antimicrobial Therapeutics Reviews

The risk/benefit of predicting a post-antibiotic era: Is the alarm working?

Tom Fowler,^{1,2} David Walker,^{3,b} and Sally C. Davies^{3,a}

¹Field Epidemiology Service–West Midlands, Public Health England, Birmingham, United Kingdom, ²Department of Public Health, Epidemiology and Biostatistics University of Birmingham, Birmingham, United Kingdom, ³Office of the Chief Medical Officer, Department of Health, London, United Kingdom

Address for correspondence: Tom Fowler, Field Epidemiology Service – West Midlands, Public Health England, Birmingham, B3 2PW, United Kingdom. Tom.Fowler@nhs.net

BAD BUGS, NO DRUGS

As Antibiotic Discovery Stagnates ...
A Public Health Crisis Brews



 **IDS**
Infectious Diseases Society of America

July 2004

Journal of Paediatrics and
Child Health



doi:10.1111/jpc.12032

VIEWPOINT

Beginning and possibly the end of the antibiotic era

Shai Ashkenazi^{1,2,3}

¹Department of Paediatrics A, Schneider Children's Medical Center of Israel, Petach Tikva; ²Sackler Faculty of Medicine and ³Felsenstein Medical Research Center, Tel Aviv University, Tel Aviv, Israel

BAD BUGS, NO DRUGS

As Antibiotic Discovery Stagnates ...
A Public Health Crisis Brews



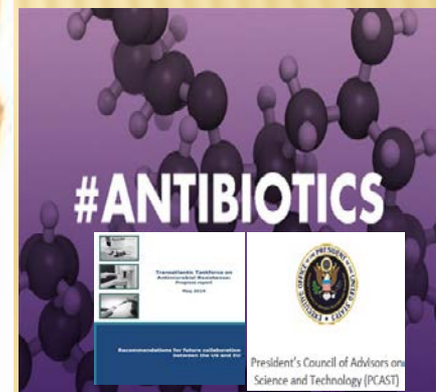
"Infectious diseases physicians are alarmed by the prospect that effective antibiotics may not be available to treat seriously ill patients in the near future. There simply aren't enough new drugs in the pharmaceutical pipeline to keep pace with drug-resistant bacterial infections, so-called 'superbugs.'"

Joseph R. Dalovisio, MD
IDSA President

OS ENORMES CUSTOS ASSOCIADOS ...

✘ Impacto da Multirresistência aos ABs (Relatórios CDC e TATFAR, 2014)

- + 20–35 bilhões USD/ano (perdas diretas)
- + 35 bilhões USD/ano (perdas de produtividade)
- + 8 milhões de dias de internamento hospitalar
- + 2 milhões de pessoas infectadas / ano
- + 23 - 99.000 mortes / ano nos EUA
- + Financiamento público na investigação de novos antibióticos insuficiente: 450 milhões USD/ano
- + Na CEE: 25 - 175.000 pessoas morrem / ano



... QUE SE REFLETE SOBRE A RIQUEZA DOS PAÍSES ...



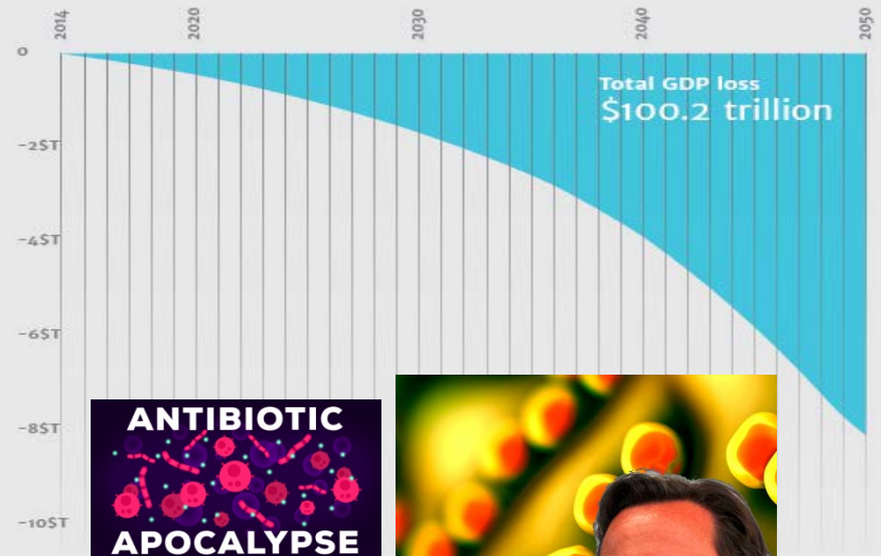
Review on
Antimicrobial
Resistance

Tackling drug-resistant infections globally

Antimicrobial
Resistance:
Tackling a crisis
for the health and
wealth of nations

The Review on Antimicrobial Resistance
Chaired by Jim O'Neill
December 2014

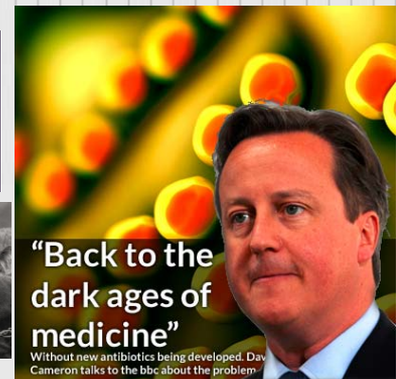
AMR's impact on World GDP
in trillions of USD



**ANTIBIOTIC
APOCALYPSE**

Antibiotic
Resistant
Bacteria are
Blowin' in
the Wind

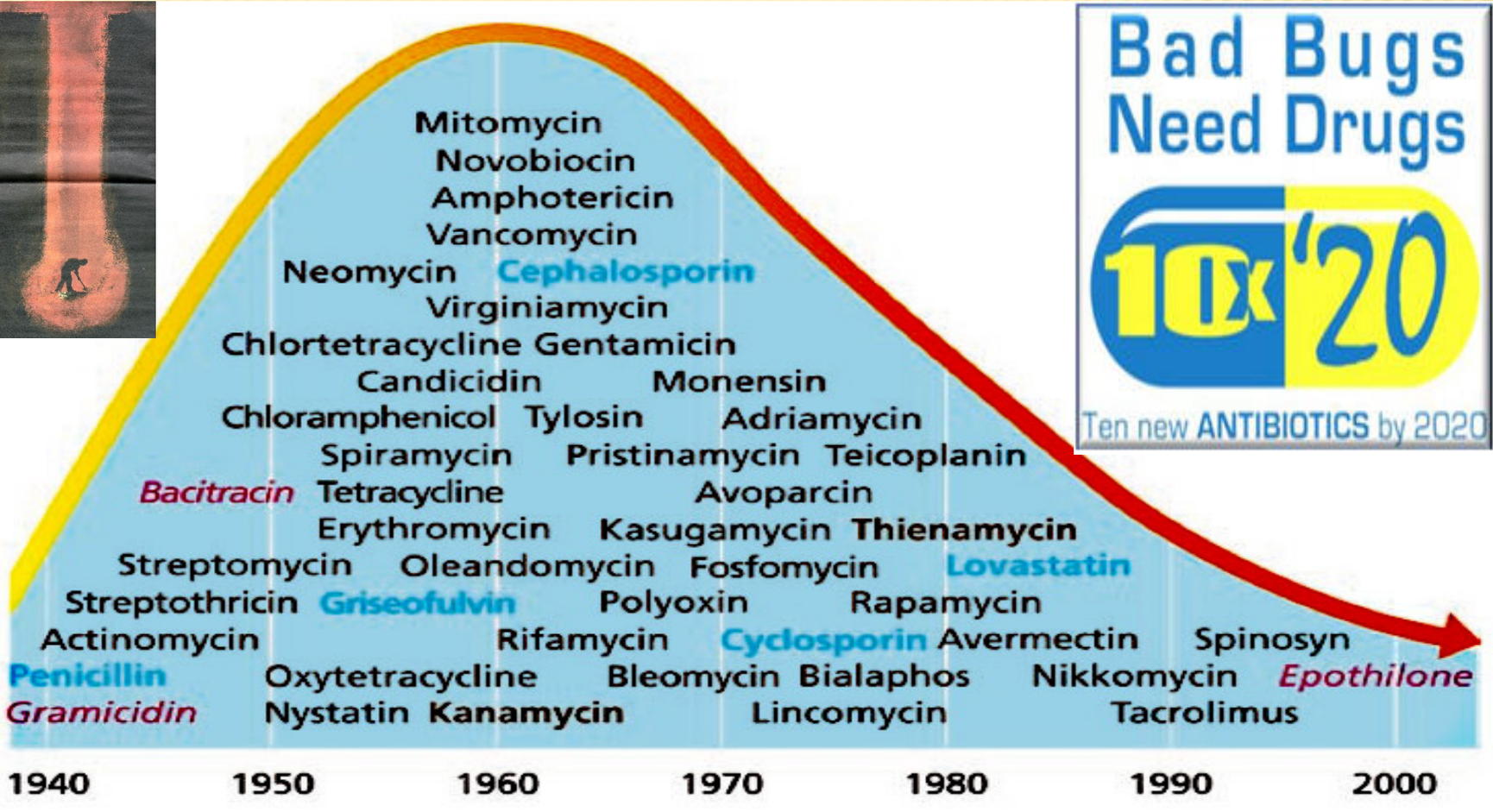
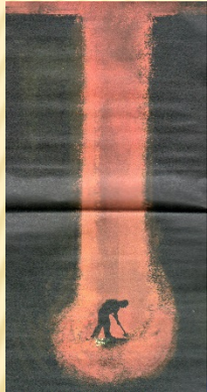
GMINSIDE



"Back to the
dark ages of
medicine"

Without new antibiotics being developed, David
Cameron talks to the BBC about the problem

A ESCASSEZ NOVOS ANTIBIÓTICOS EM DESENVOLVIMENTO ATÉ AO FIM DO SEC. XX...



... QUE NÃO FOI RESOLVIDA NA PRIMEIRA DÉCADA DO SEC. XXI ...

The Antibiotic Resistance Crisis

Part 1: Causes and Threats

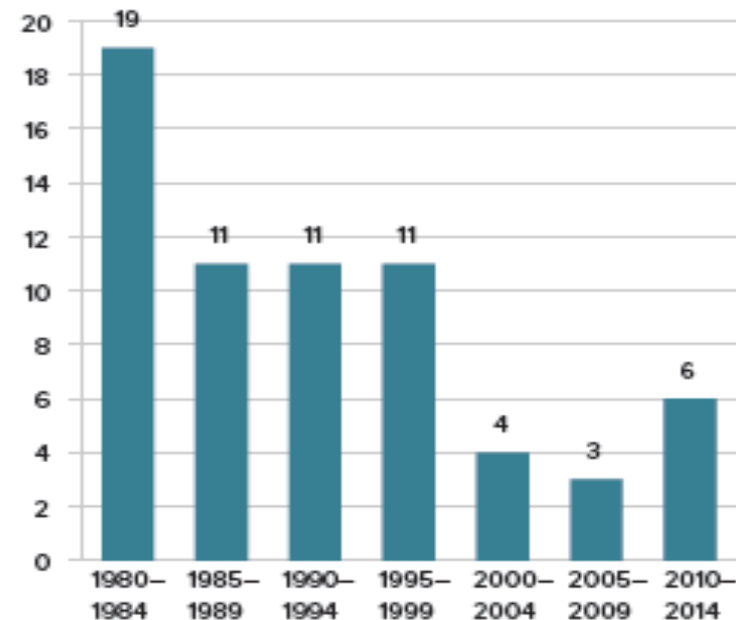
C. Lee Ventola, MS

The Antibiotic Resistance Crisis

Part 2: Management Strategies and New Agents

C. Lee Ventola, MS

Figure 3 Number of Antibacterial New Drug Application Approvals Versus Year Intervals



The number of new antibiotics developed and approved has decreased steadily over the past three decades (although four new drugs were approved in 2014), leaving fewer options to treat resistant bacteria.

* Drugs are limited to systemic agents. Data courtesy of the CDC⁵ and the FDA Center for Drug Evaluation and Research.

... QUE NÃO PERMITE RESPONDER AO PROBLEMA DAS BACTÉRIAS MULTIRESISTENTES ...

Figure 1 Developing Antibiotic Resistance: A Timeline of Key Events⁵

ANTIBIOTIC RESISTANCE IDENTIFIED	ANTIBIOTIC INTRODUCED
Penicillin-R <i>Staphylococcus</i> 1940	1943 Penicillin
	1950 Tetracycline
	1953 Erythromycin
Tetracycline-R <i>Shigella</i> 1959	1960 Methicillin
Methicillin-R <i>Staphylococcus</i> 1962	1967 Gentamicin
Penicillin-R pneumococcus 1965	1972 Vancomycin
Erythromycin-R <i>Streptococcus</i> 1968	
	1985 Imipenem and ceftazidime
Gentamicin-R <i>Enterococcus</i> 1979	
Ceftazidime-R Enterobacteriaceae 1987	
Vancomycin-R <i>Enterococcus</i> 1988	
	1996 Levofloxacin
Levofloxacin-R pneumococcus 1996	
Imipenem-R Enterobacteriaceae 1998	2000 Linezolid
XDR tuberculosis 2000	
Linezolid-R <i>Staphylococcus</i> 2001	
Vancomycin-R <i>Staphylococcus</i> 2002	2003 Daptomycin
PDR- <i>Acinetobacter</i> and <i>Pseudomonas</i> 2004/5	
	2010 Ceftaroline
Ceftriaxone-R <i>Neisseria gonorrhoeae</i> 2009	
PDR-Enterobacteriaceae 2009	
Ceftaroline-R <i>Staphylococcus</i> 2011	

PDR = pan-drug-resistant; R = resistant; XDR = extensively drug-resistant
 Dates are based upon early reports of resistance in the literature. In the case of pan-drug-resistant *Acinetobacter* and *Pseudomonas*, the date is based upon reports of health care transmission or outbreaks. Note: penicillin was in limited use prior to widespread population usage in 1943.

Table 1 CDC Assessment of Antibacterial Resistance Threats⁵

Urgent Threats

- *Clostridium difficile*
- Carbapenem-resistant Enterobacteriaceae (CRE)
- Drug-resistant *Neisseria gonorrhoeae*

Serious Threats

- Multidrug-resistant *Acinetobacter*
- Drug-resistant *Campylobacter*
- Fluconazole-resistant *Candida* (a fungus)
- Extended spectrum beta-lactamase-producing Enterobacteriaceae (ESBLs)
- Vancomycin-resistant Enterococci (VRE)
- Multidrug-resistant *Pseudomonas aeruginosa*
- Drug-resistant nontyphoidal *Salmonella*
- Drug-resistant *Salmonella* Typhimurium
- Drug-resistant *Shigella*
- Methicillin-resistant *Staphylococcus aureus* (MRSA)
- Drug-resistant *Streptococcus pneumoniae*
- Drug-resistant tuberculosis

Concerning Threats

- Vancomycin-resistant *Staphylococcus aureus* (VRSA)
- Erythromycin-resistant Group A *Streptococcus*
- Clindamycin-resistant Group B *Streptococcus*

... A RESPOSTA NO 2ª DÉCADA DO SEC XXI

The Antibiotic Resistance Crisis, Part 2: Management Strategies and New Agents

Table 1 CDC's Antibiotic Resistance and Antibiotic-Resistant Infections Tracking Platform ⁵		
Tracking Networks	Data Collected	Resistant Bacteria/Fungi*
Emerging Infections Program (EIP) , which has three main elements: <ul style="list-style-type: none"> • ABCs: Active Bacterial Core surveillance • HAIC: Healthcare-Associated Infections—Community Interface • FoodNet: Foodborne Diseases Active Surveillance Network 	A network of public health/academic/hospital collaborations in 10 states. It provides access to bacterial and fungal samples for testing and detailed clinical case data. The three main EIP programs collect different types of resistance data: <ul style="list-style-type: none"> • ABCs provides clinical information and resistance data for bacteria that cause infections predominantly in the community. • HAIC provides clinical information and resistance data for bacteria and fungi that cause infections at the intersection of health care and the general community. • FoodNet supplies clinical and epidemiologic data on some human isolates in the National Antimicrobial Resistance Monitoring System (NARMS). 	ABCs <ul style="list-style-type: none"> • <i>Streptococcus pneumoniae</i> • Groups A and B <i>Streptococcus</i> • Methicillin-resistant <i>Staphylococcus aureus</i> HAIC <ul style="list-style-type: none"> • <i>Clostridium difficile</i> • <i>Candida</i> (a fungus) • Carbapenem-resistant Enterobacteriaceae • Multiple-drug-resistant <i>Acinetobacter</i> FoodNet: (see NARMS list below)
National Antimicrobial Resistance Monitoring System (NARMS)	A national public health surveillance system that tracks changes in the susceptibility of foodborne and other enteric bacteria to antibiotics of human and veterinary medical importance. NARMS is a collaboration among the CDC, FDA, USDA, and state and local health departments. CDC tests bacterial isolates from humans, while FDA and USDA test isolates from retail meats and food animals.	<ul style="list-style-type: none"> • <i>Salmonella</i> • <i>Campylobacter</i> • <i>Shigella</i>
National Healthcare Safety Network (NHSN)	A system that collects and provides data on infections and drug resistance in health care settings. Since NHSN collects data directly from health care facilities, it can provide facility-level information on health care-associated infections and antibiotic resistance (and, in the future, on antibiotic use).	<ul style="list-style-type: none"> • <i>S. aureus</i> • <i>Enterococcus</i> • Enterobacteriaceae • <i>Acinetobacter</i> • <i>Pseudomonas aeruginosa</i> • <i>Candida</i>
Gonococcal Isolate Surveillance Program (GISP)	A program to track antibiotic resistance data for gonococcal isolates. Isolates are collected from sexually transmitted disease clinics in approximately 28 cities.	<i>Neisseria gonorrhoeae</i>
National Tuberculosis Surveillance System (NTSS)	National Electronic Disease Surveillance System–based reporting of tuberculosis cases, including resistance data. Public health departments from 50 states and the U.S. territories contribute data.	<i>Mycobacterium tuberculosis</i>

* ABCs also includes surveillance for *Neisseria meningitidis* and *Haemophilus influenzae*. NARMS also includes surveillance for *E. coli O157* and *Vibrio* (non-*V. cholerae*).

CDC = Centers for Disease Control and Prevention; FDA = Food and Drug Administration; USDA = U.S. Department of Agriculture

The Antibiotic Resistance Crisis, Part 2: Management Strategies and New Agents

Table 2 Antibiotics Approved Since 2005 ^{5,20–23}			
Drug Name	Year Approved	Drug Class	Indications*
Tigecycline (Tygacil, Pfizer)	2005	Tetracycline	Patients 18 years of age and older with: <ul style="list-style-type: none"> • cSSSIs • cAIs • CABP
Doripenem (Doribax, Shionogi)	2007	Carbapenem	Adult patients with: <ul style="list-style-type: none"> • cAIs • cUTIs, including pyelonephritis
Telavancin (Vibativ, Theravance Biopharma)	2008	Glycopeptide	Adult patients with: <ul style="list-style-type: none"> • Hospital-acquired and ventilator-associated bacterial pneumonia • cSSSIs
Ceftaroline (Teflaro, Cerexa)	2010	Cephalosporin	Patients with:† <ul style="list-style-type: none"> • CABP • ABSSSIs
Tedizolid (Sivextro, Cubist Pharmaceuticals)	2014	Oxazolidinone	Adult patients with ABSSSIs
Dalbavancin (Dalvance, Durata Therapeutics)	2014	Glycopeptide	Adult patients with ABSSSIs
Oritavancin (Orbactiv, Medicines Company)	2014	Glycopeptide	Adult patients with ABSSSIs
Ceftolozane/tazobactam (Zerbaxa, Cubist Pharmaceuticals)	2014	Cephalosporin/beta-lactamase inhibitor	Adult patients with: <ul style="list-style-type: none"> • cAIs, in combination with metronidazole • cUTIs, including pyelonephritis
Ceftazidime/avibactam (Avycaz, Cerexa Inc.)	2015	Cephalosporin/beta-lactamase inhibitor	Adult patients with: <ul style="list-style-type: none"> • cAIs, in combination with metronidazole • cUTIs, including pyelonephritis

ABSSSIs = acute bacterial skin and skin structure infections; CABP = community-acquired bacterial pneumonia; cAIs = complicated intra-abdominal infections; cSSSIs = complicated skin and skin structure infections; cUTIs = complicated urinary tract infections

* Caused by designated susceptible bacteria. See prescribing information for these and other important details.

† Safety and effectiveness in pediatric patients have not been established.

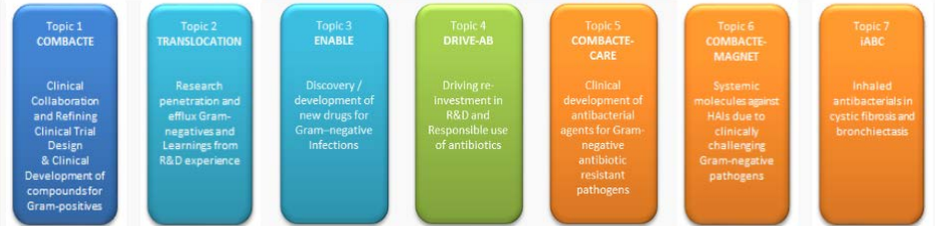
ATRAVÉS DA COOPERAÇÃO ENTRE DIVERSOS ORGANISMOS OFICIAIS E PRIVADOS A NÍVEL INTERNACIONAL ...

The right prevention and treatment
for the right patient at the right time

Strategic Research Agenda for
Innovative Medicines Initiative 2



Structure of New Drugs 4 Bad Bugs (ND4BB)



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... COMO NO CASO DO TETFAR (USA/CEE)



Transatlantic Taskforce on
Antimicrobial Resistance:
Progress report

May 2014



Recommendations for future collaboration
between the US and EU

WASHINGTON JOURNAL OF LAW, TECHNOLOGY & ARTS
VOLUME 9, ISSUE 1 SUMMER 2013

REPAIRING THE ANTIBIOTIC PIPELINE: CAN THE GAIN ACT DO IT?

Caitlin Forsyth¹
© Caitlin Forsyth

Cite as: 9 WASH. J.L. TECH. & ARTS 1 (2013)
<https://digital.lib.washington.edu/dspace-law/handle/1773.1/1267>

ABSTRACT

Antibiotic resistance, according to the World Health Organization, is one of the greatest threats to public health. To combat the problem, new antibiotics need to be developed. However, antibiotic research and development is fraught with scientific and economic problems. Recognizing these problems and the public health threat posed by antibiotic resistance, Congress passed the GAIN Act, which President Obama signed into law in June 2012. The GAIN Act (Act) incentivizes pharmaceutical companies to invest in antibiotic research and development. This Article will outline the incentives in the Act and suggest why the Act may not solve the growing antibiotic resistance problem. There are, however, areas of promise in the Act that may mitigate its shortcomings and pave the way to the possibility of the Act's success.



REPORT TO THE PRESIDENT ON COMBATING ANTIBIOTIC RESISTANCE

Executive Office of the President
President's Council of Advisors on
Science and Technology

September 2014



OS "TIMELINES" QUE IMPORTA TER CONSIDERAR !!!

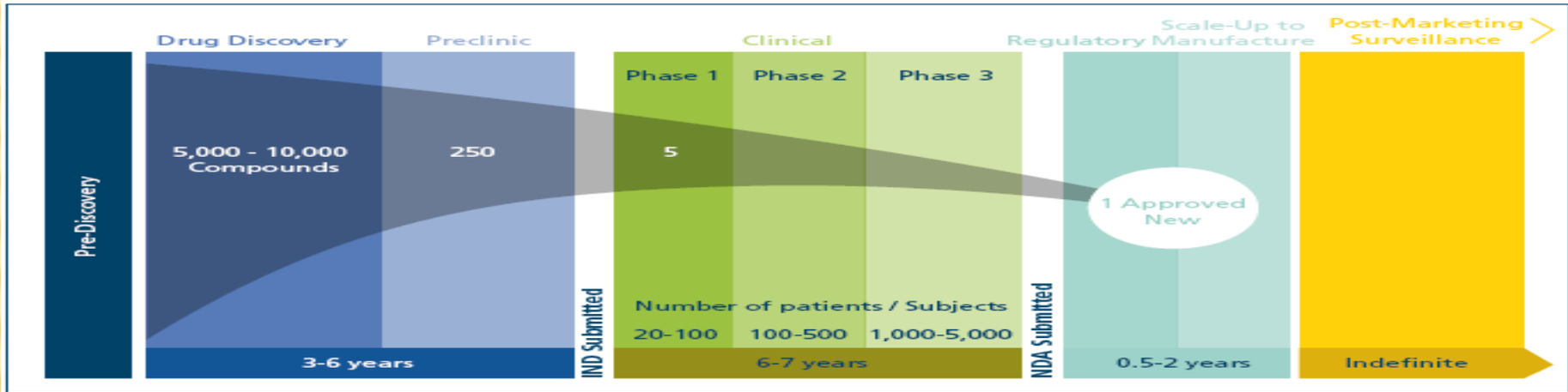


Figure 6: With the cost of developing a medicine estimated at \$1.2 billion, attrition rates in pharmaceutical R&D are unsustainable. Source CBO, Research and Development in the Pharmaceutical Industry, 2006.

Annex E: Timeline of the Transatlantic Task Force on Antimicrobial Resistance (TATFAR), 2009-2013

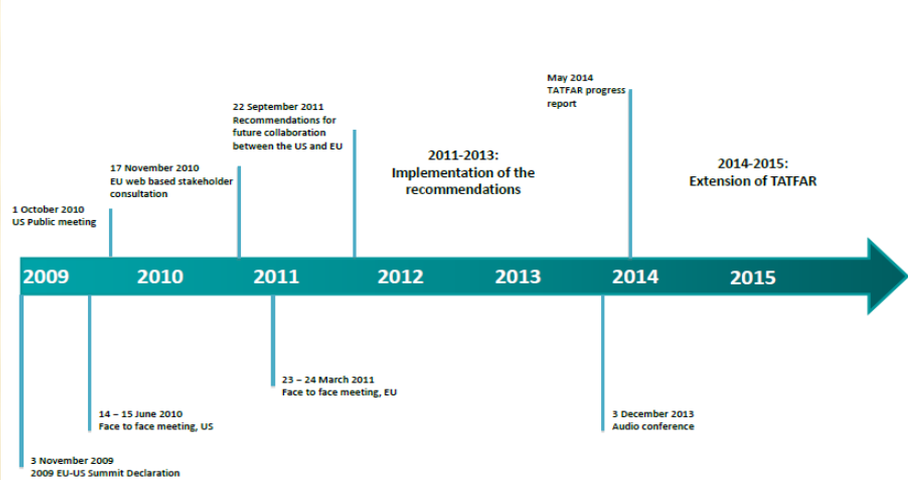
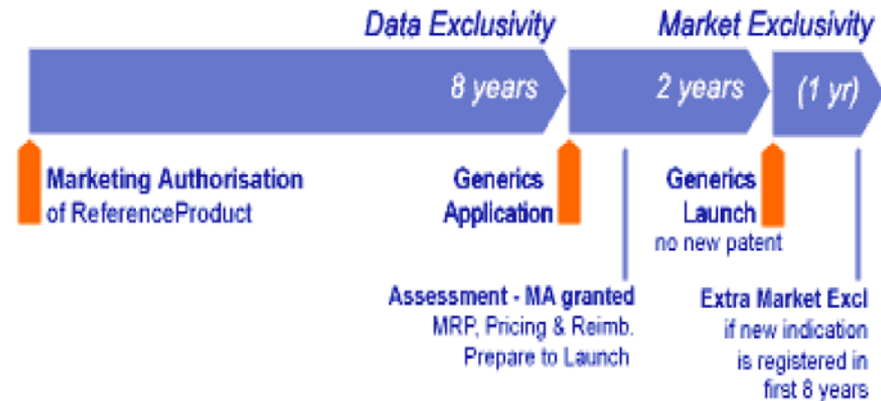
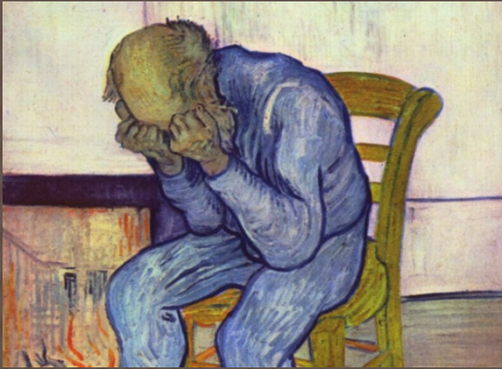


Figure 6.2.1 EMEA 8+2(+1) arrangements in the EU



É BOM SEMPRE LEMBRAR!!!

- ✘ “Guidelines” internacionais recomendam
 - + Sinusite Aguda: esperar 1 semana antes de iniciar ABs
 - + Bronquite aguda: não deve ser TT de início c/ ABs
 - + Faringite aguda: c/ Penicilina G e apenas nos casos c/ Ag + p/ o *Streptococcus Pyogenes*
 - + Síndromes gripais e as constipações não devem ser tratadas de início c/ ABs
 - + ITUs assintomáticas também não devem ser tratadas c/ ABs



III)- AS DOENÇAS EMERGENTES
(E REEMERGENTES): AS
INFEÇÕES TRANSMITIDAS POR
VETORES E ZOOLOSES

A CONSTATAÇÃO DE QUE TODOS OS FATORES ENVOLVIDOS ESTÃO INTERLIGADOS ...

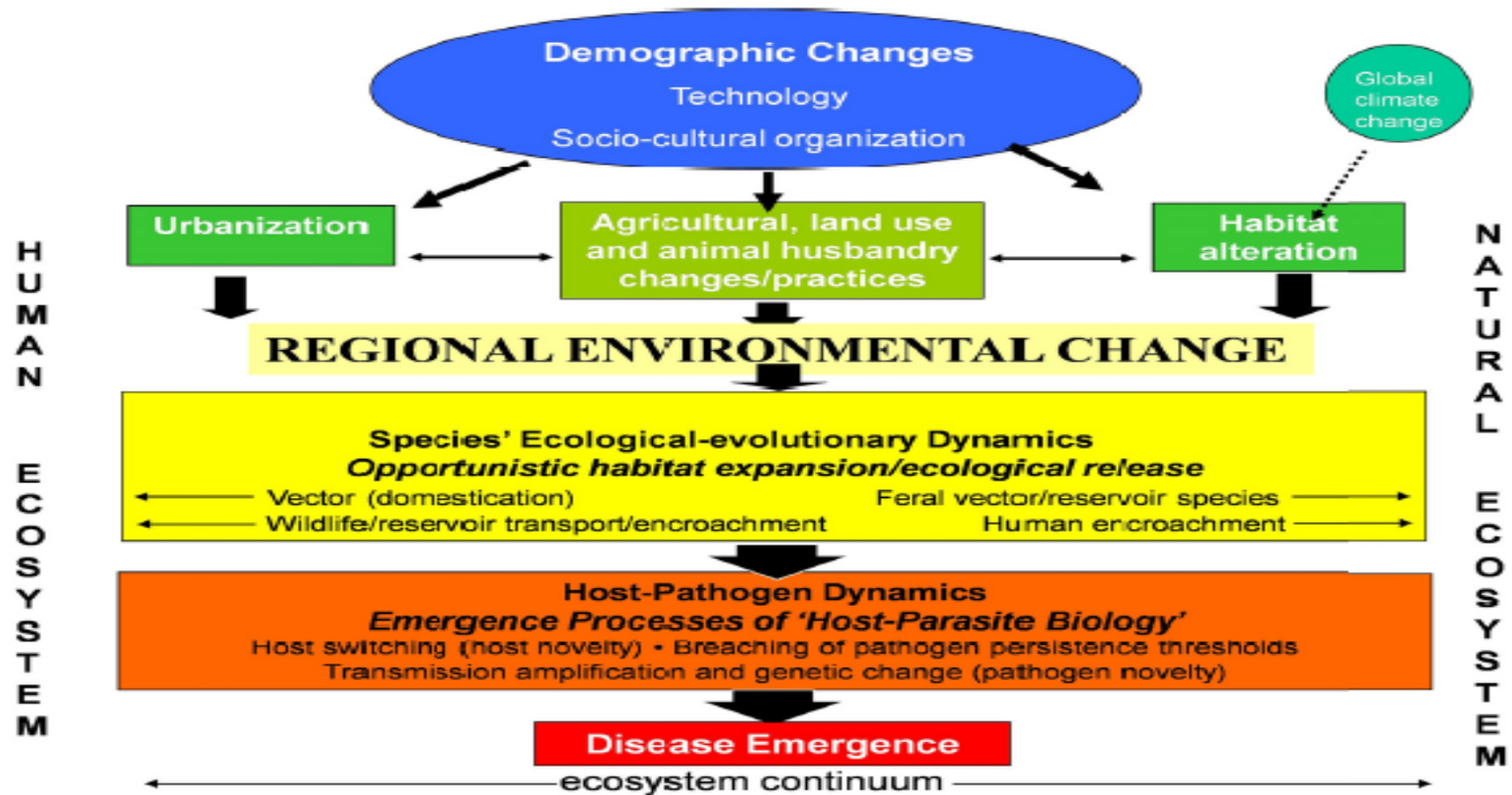


FIGURE WO-5 Epidemiological effects of urbanization and environmental change.
SOURCE: Adapted from Wilcox and Gubler (2005) with permission from The Japanese Society for Hygiene.

... SOBRETUDO NO QUE SE REFERE ÀS DOENÇAS EMERGENTES ...

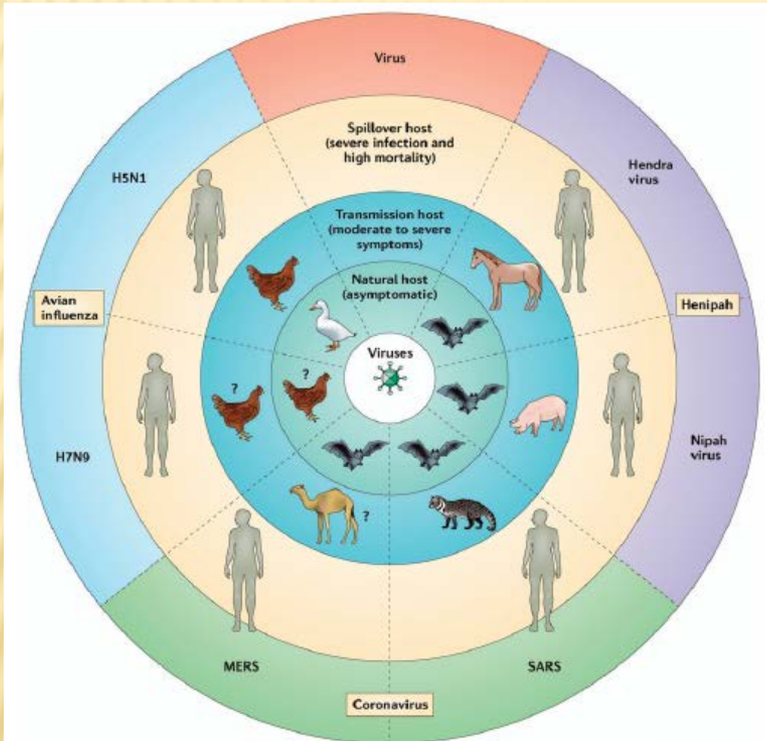


FIGURE A6-2 The outcome of disease severity is influenced by the host–pathogen interaction. Many zoonotic agents cause little or no signs of disease in their natural host such as wild birds and bats, while transmission hosts may present symptoms ranging from moderate (such as pigs for AI) to severe (such as horses for HeV) signs. The terminal or spillover host, such as humans in the case of H5N1 and HeV infections, can present with very severe symptoms and high mortality rates. For some of the most recently emerging EIDs such as H7N9 and MERS-CoV, natural and transmission hosts have not been identified. SOURCE: Bean et al., 2013.

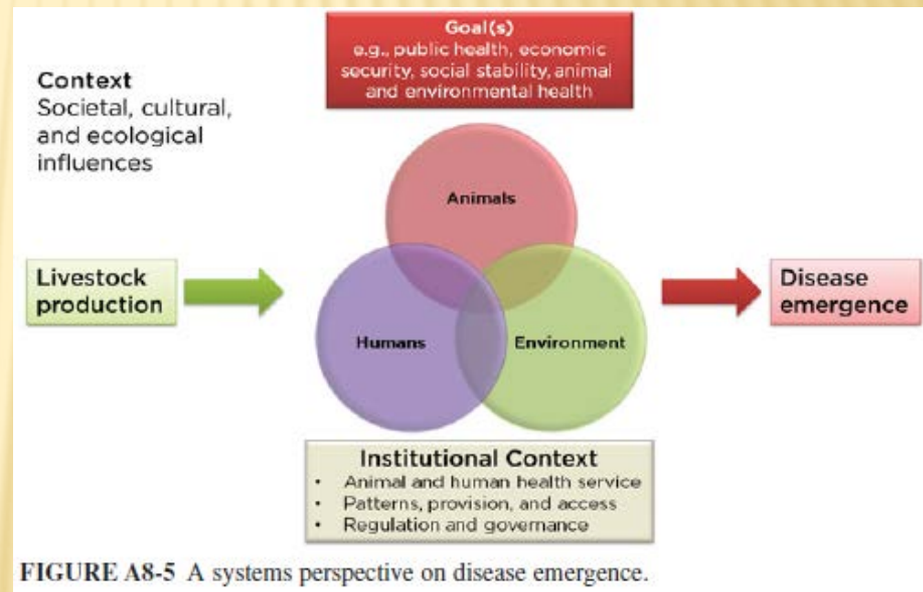


FIGURE A8-5 A systems perspective on disease emergence.

... E PARTICULARMENTE NO CASOS DAS INFECÇÕES VIRAIS ...

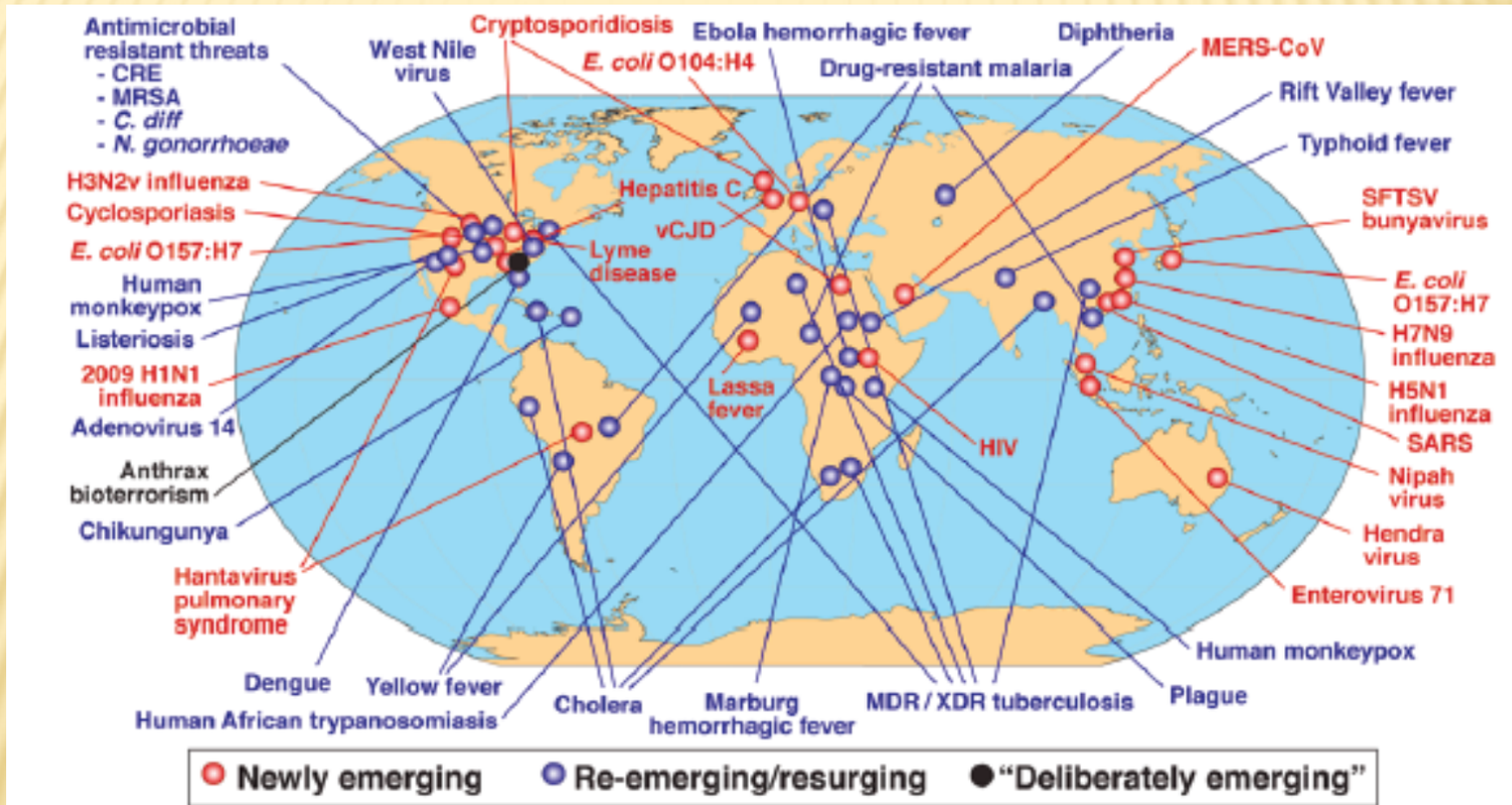


FIGURE WO-1 Global examples of emerging and reemerging infectious diseases.
 SOURCE: Morens et al., 2004.

... QUE APRESENTARAM, AO LONGO DO TEMPO, UM GRANDE IMPACTO EPIDEMIOLÓGICO!!!

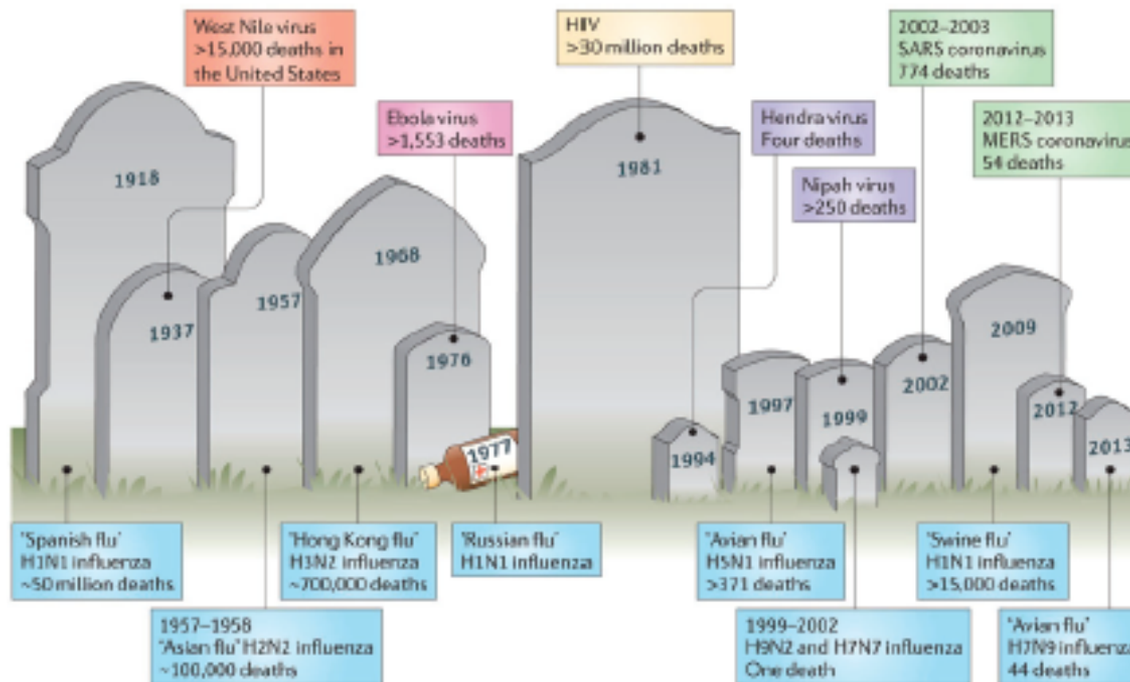


FIGURE WO-2 Emergence of zoonoses. Over the past century, humanity has witnessed the emergence of numerous zoonotic infections that have resulted in varying numbers of human fatalities. Influenza viruses that originate from birds account for an important proportion of these deaths, and recently many new zoonotic viruses that originate in bats, such as Hendra virus, Nipah virus, and the SARS coronavirus, have caused outbreaks with high mortality rates.

NOTE: As of June 2, 2014, the Centers for Disease Control and Prevention (CDC) reports that there were 39,557 cases of West Nile virus in the United States resulting in 1,668 deaths between 1999 and 2013. Source: http://www.cdc.gov/westnile/resources/pdfs/cummulative/99_2013_CasesAndDeathsClinicalPresentationHumanCases.pdf (accessed February 19, 2015).

SOURCE: Bean et al., 2013.

O ENORME IMPACTO ECONÓMICO DAS INFEÇÕES EMERGENTES

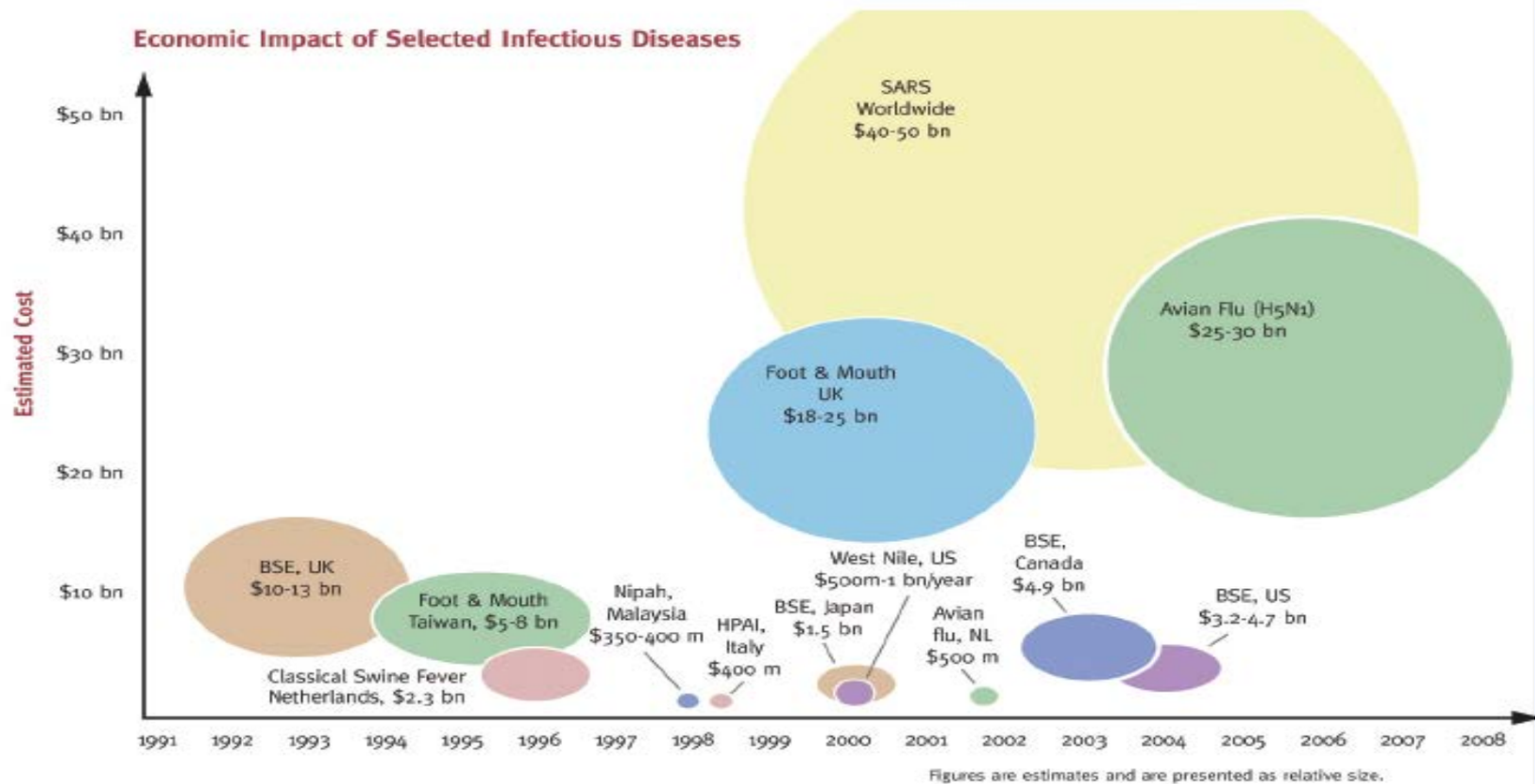
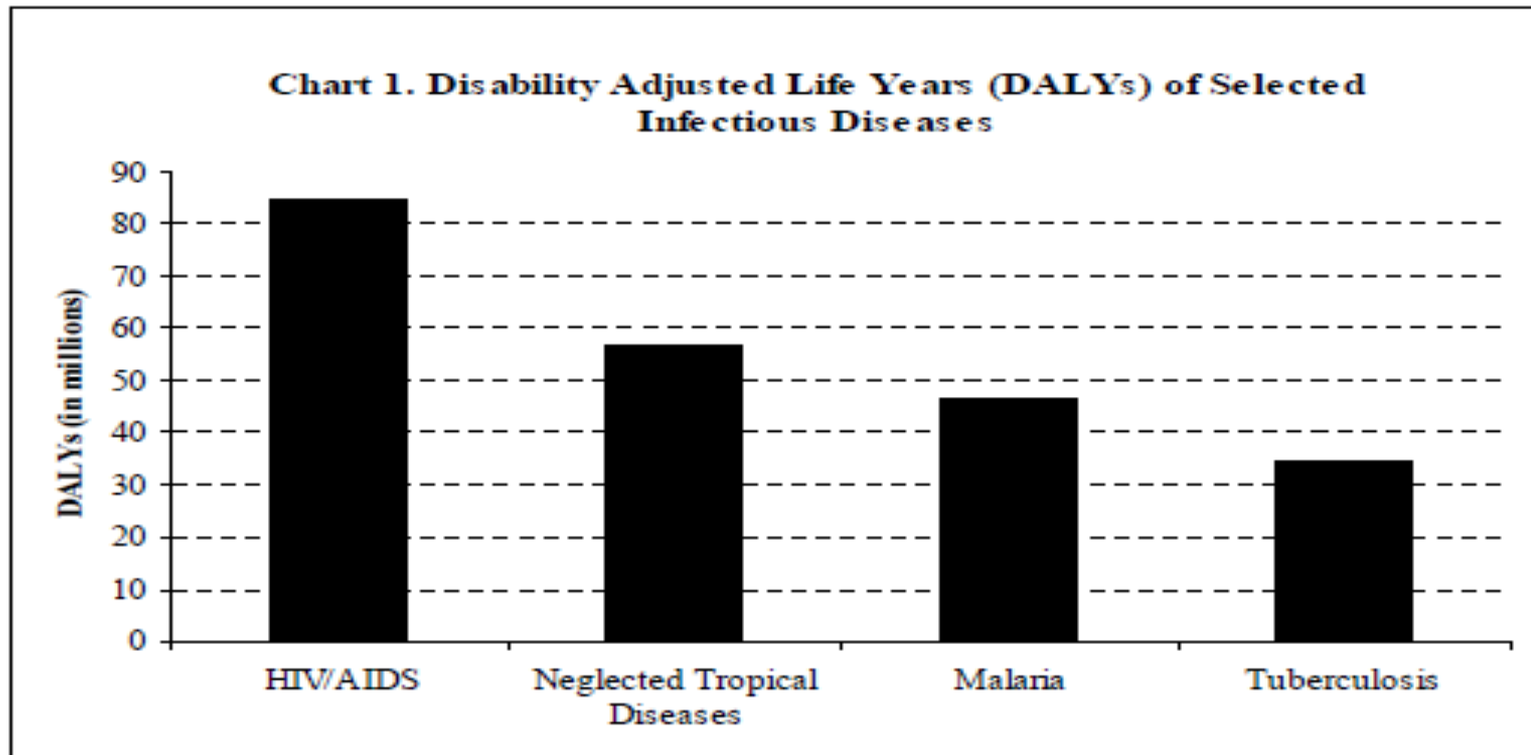


FIGURE 3-2 Economic impacts of selected infectious diseases.
SOURCE: Reprinted with permission from Bio-era.

A IMPORTÂNCIA DAS DENOMINADAS DOENÇAS NEGLIGENCIADAS ...

The unquestionable association between NTDs and poverty has made their elimination a necessity in order to achieve the Millennium Development Goals (MDGs).



* Data taken Hotez PJ, Molyneux DH, Fenwick A et al. Control of Neglected Tropical Diseases. N Engl J Med 2007; 357:1018-1027.

... A SUA DISTRIBUIÇÃO GEOGRÁFICA MUNDIAL

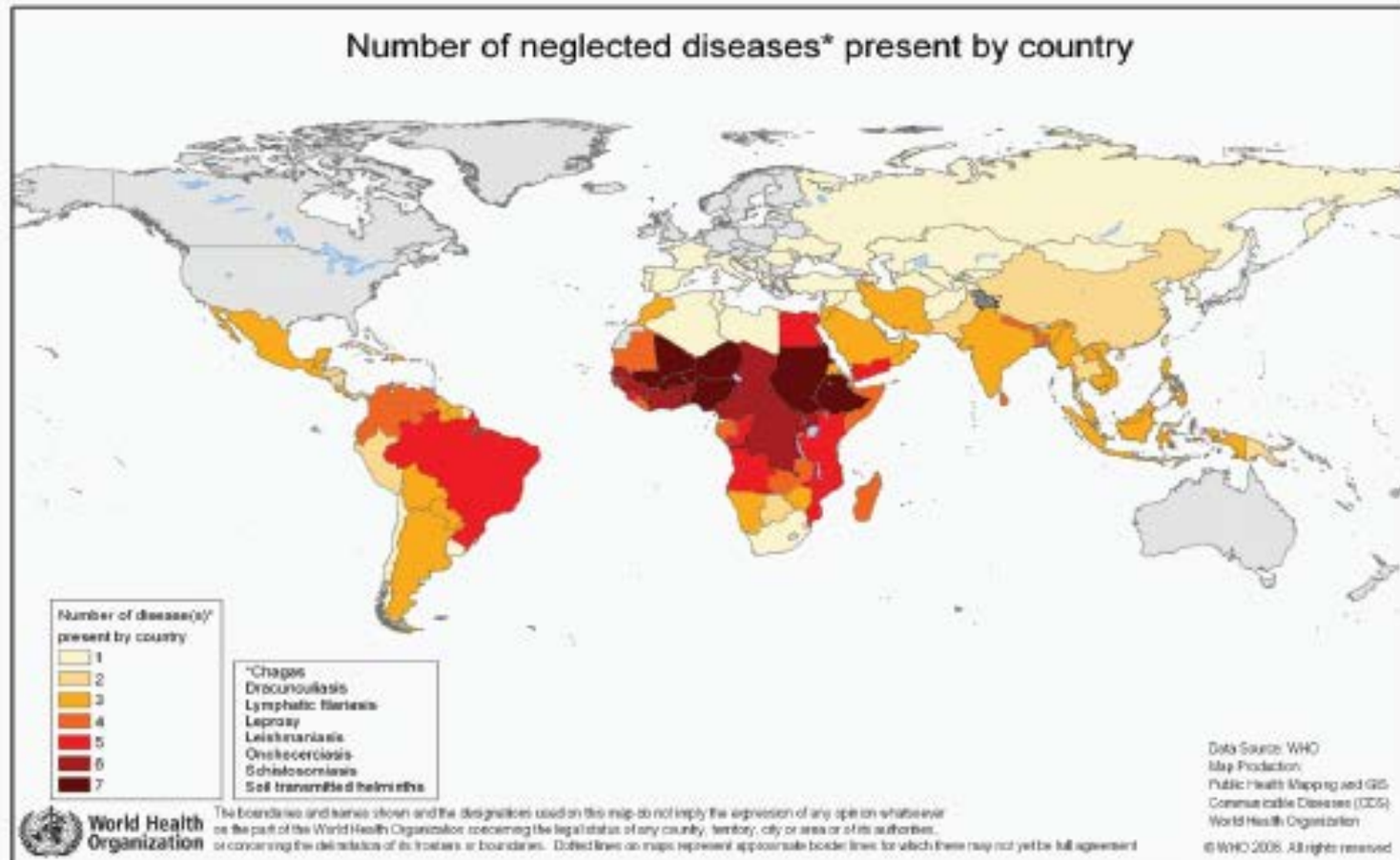


FIGURE A15-4 Global distribution of NTDs.

SOURCE: WHO (2006). *Number of neglected diseases present by country* [map]. Geneva: WHO. Map produced by Public Health Mapping and GIS, Communicable Diseases, WHO (2006). Reprinted with permission from the World Health Organization.

NOVOS MOTIVOS DE PREOCUPAÇÃO!!!

Review

J Vet Sci 2016, 17(1), 1-11 • <http://dx.doi.org/10.4142/jvs.2016.17.1.1>

JVS

Hepatitis E virus as an emerging zoonotic pathogen

Woo-Jung Park¹, Byung-Joo Park¹, Hee-Seop Ahn¹, Joong-Bok Lee¹, Seung-Yong Park¹, Chang-Seon Song¹, Sang-Won Lee¹, Han-Sang Yoo², In-Soo Choi^{1,*}

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Hepatitis E outbreaks are a serious public health concern in developing countries. The disease causes acute infections, primarily in young adults. The mortality rate is approximately 2%; however, it can exceed 20% in pregnant women in some regions in India. The causative agent, hepatitis E virus (HEV), has been isolated from several animal species, including pigs. HEV genotypes 3 and 4 have been isolated from both humans and animals, and are recognized as zoonotic pathogens. Seroprevalence studies in animals and humans indirectly suggest that HEV infections occur worldwide. The virus is primarily transmitted to humans via undercooked animal meats in developed countries. Moreover, transfusion- and transplantation-mediated HEV infections have recently been reported. This review summarizes the general characteristics of hepatitis E, HEV infection status in animals and humans, the zoonotic transmission modes of HEV, and HEV vaccine development status.

Keywords: hepatitis E, hepatitis E virus, pig, transmission, zoonotic pathogen

NOVAS REALIDADES EPIDEMIOLÓGICAS PODEM FAZER DA EXCEÇÃO A REGRA

Furuya-Kanamori et al. *BMC Infectious Diseases* (2016) 16:84
DOI 10.1186/s12879-016-1417-2

BMC Infectious Diseases

RESEARCH ARTICLE

Open Access

Co-distribution and co-infection of chikungunya and dengue viruses

Luis Furuya-Kanamori^{1*}, Shaohong Liang², Gabriel Milinovich³, Ricardo J. Soares Magalhaes^{4,5},
Archie C. A. Clements¹, Wenbiao Hu³, Patricia Brasil⁶, Francesca D. Frentiu⁷, Rebecca Dunning⁸ and Laith Yakob⁹



Abstract

Background: Chikungunya and dengue infections are spatio-temporally related. The current review aims to determine the geographic limits of chikungunya, dengue and the principal mosquito vectors for both viruses and to synthesise current epidemiological understanding of their co-distribution.

Methods: Three biomedical databases (PubMed, Scopus and Web of Science) were searched from their inception until May 2015 for studies that reported concurrent detection of chikungunya and dengue viruses in the same patient. Additionally, data from WHO, CDC and Healthmap alerts were extracted to create up-to-date global distribution maps for both dengue and chikungunya.

Results: Evidence for chikungunya-dengue co-infection has been found in Angola, Gabon, India, Madagascar, Malaysia, Myanmar, Nigeria, Saint Martin, Singapore, Sri Lanka, Tanzania, Thailand and Yemen; these constitute only 13 out of the 98 countries/territories where both chikungunya and dengue epidemic/endemic transmission have been reported.

Conclusions: Understanding the true extent of chikungunya-dengue co-infection is hampered by current diagnosis largely based on their similar symptoms. Heightened awareness of chikungunya among the public and public health practitioners in the advent of the ongoing outbreak in the Americas can be expected to improve diagnostic rigour. Maps generated from the newly compiled lists of the geographic distribution of both pathogens and vectors represent the current geographical limits of chikungunya and dengue, as well as the countries/territories at risk of future incursion by both viruses. These describe regions of co-endemicity in which lab-based diagnosis of suspected cases is of higher priority.

Keywords: Chikungunya, Dengue, Virus, Coinfection, Review

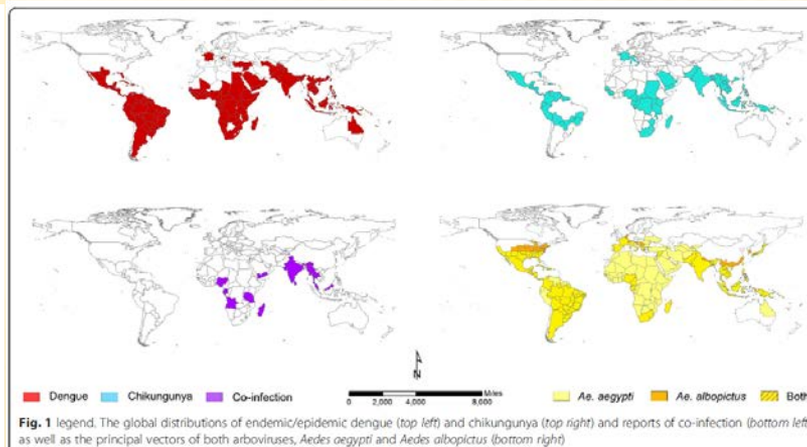


Fig. 1 legend. The global distributions of endemic/epidemic dengue (top left) and chikungunya (top right) and reports of co-infection (bottom left) as well as the principal vectors of both arboviruses, *Aedes aegypti* and *Aedes albopictus* (bottom right)

RAPID COMMUNICATIONS

Experimental studies of susceptibility of Italian *Aedes albopictus* to Zika virus

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AS DOENÇAS DA POBREZA ATINGEM TAMBÉM OS PAÍSES DENOMINADOS DE RICOS...

Zhou et al. *Infectious Diseases of Poverty* (2016) 5:49
DOI 10.1186/s40249-016-0144-7

Infectious Diseases of Poverty

COMMENTARY

Open Access

Surveillance and response systems for elimination of tropical diseases: summary of a thematic series in *Infectious Diseases of Poverty*



Xia Zhou^{1,2}, Peiling Yap^{3,4}, Marcel Tanner^{3,4}, Robert Bergquist⁵, Jürg Utzinger^{3,4} and Xiao-Nong Zhou^{2,6*}

Abstract

The peer-reviewed journal *Infectious Diseases of Poverty* provides a new platform to engage with, and disseminate in an open-access format, science outside traditional disciplinary boundaries. The current piece reviews a thematic series on surveillance-response systems for elimination of tropical diseases. Overall, 22 contributions covering a broad array of diseases are featured – i.e. clonorchiasis, dengue, hepatitis, human immunodeficiency virus/acquired immune deficiency syndrome (HIV/AIDS), H7N9 avian influenza, lymphatic filariasis, malaria, Middle East respiratory syndrome (MERS), rabies, schistosomiasis and tuberculosis (TB). There are five scoping reviews, a commentary, a letter to the editor, an opinion piece and an editorial pertaining to the theme “Elimination of tropical disease through surveillance and response”. The remaining 13 articles are original contributions mainly covering (i) drug resistance; (ii) innovation and validation in the field of mathematical modelling; (iii) elimination of infectious diseases; and (iv) social media reports on disease outbreak notifications released by national health authorities. Analysis of the authors’ affiliations reveals that scientists from the People’s Republic of China (P.R. China) are prominently represented. Possible explanations include the fact that the 2012 and 2014 international conferences pertaining to surveillance-response mechanisms were both hosted by the National Institute of Parasitic Diseases (NIPD) in Shanghai, coupled with P.R. China’s growing importance with regard to the control of infectious diseases. Within 4 to 22 months of publication, three of the 22 contributions were viewed more than 10 000 times each. With sustained efforts focusing on relevant and strategic information towards control and elimination of infectious diseases, *Infectious Diseases of Poverty* has become a leading journal in the field of surveillance and response systems in infectious diseases and beyond.

Keywords: Infectious diseases, Tropical diseases, Health systems, Surveillance and response systems, Elimination, People’s Republic of China

Journal of Epidemiology and Global Health (2016) xxx, xxx–xxx



FI SEVIER

<http://www.elsevier.com/locate/jegh>



REVIEW ARTICLE

Bradford Hill’s criteria, emerging zoonoses, and One Health

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Received 1 September 2015; accepted 19 October 2015

KEYWORDS

Disease causation;
Emerging zoonoses;
One Health

Abstract Zoonoses constitute more than 60% of infectious diseases and 75% of emerging infectious diseases. Inappropriate overemphasis of specialization of disciplines has ignored public health. Identifying the causes of disease and determining how exposures are related to outcomes in “emerging zoonoses” affecting multiple species are considered to be the hallmarks of public health research and practice that compels the adoption of “One Health”. The interactions within and among populations of vertebrates in the causation and transmissions of emerging zoonotic diseases are inherently dynamic, interdependent, and systems based. Disease causality theories have moved from one or several agents causing disease in a single species, to one infectious agent causing disease in multiple species-emerging zoonoses. Identification of the causative pathogen components or structures, elucidating the mechanisms of species specificity, and understanding the natural conditions of emergence would facilitate better derivation of the causal mechanism. Good quality evidence on causation in emerging zoonoses affecting multiple species makes a strong recommendation under the One Health approach for disease prevention and control from diagnostic tests, treatment, antimicrobial resistance, preventive vaccines, and evidence informed health policies. In the tenets of One Health, alliances work best when the legitimate interests of the different partners combine to prevent and control emerging zoonoses.

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... REALIDADE A QUE A VELHA EUROPA NÃO ESCAPA...

PERSPECTIVE

Determinants and Drivers of Infectious Disease Threat Events in Europe

Jan C. Semenza, Elisabet Lindgren, Laszlo Balkanyi, Laura Espinosa, My S. Almqvist, Pasi Penttinen, Joacim Rocklöv

Table 2. Infectious disease threat events detected in Europe, 2008–2013

Threat event category	Definition and examples ^a
Foodborne and waterborne	All types of diseases caused by the transmission of organisms through food or water (e.g., drinking water, recreational water): salmonellosis, hepatitis A, <i>Escherichia coli</i> infection, norovirus infection, shigellosis.
Vectorborne and rodentborne	All vectorborne and rodentborne diseases (epidemics or first autochthonous cases): West Nile fever, malaria, dengue fever, Hantavirus infection.
Other zoonoses	Diseases caused by transmission of organisms through contact with animals or animal discharges: Q fever, cowpox disease, psittacosis.
Vaccine preventable	Main vaccine-preventable diseases that are normally part of the public health system's vaccination programs: measles, pertussis, mumps (boys), rubella (girls).
Multidrug resistance associated	Emerging multidrug-resistant infections of public health concern: carbapenemase-producing <i>Enterobacteriaceae</i> , <i>Klebsiella pneumoniae</i> .
Healthcare associated	Infections contracted while hospitalized or transmitted through healthcare practices: meningococcal meningitis.
Injection drug use associated	Infections caused by injection drug use: botulism, HIV, anthrax.
Sexually transmitted	Emerging sexually transmitted diseases and increases in incidence of serious complications: meningococcal infections.
Influenza	Seasonal influenza and other pandemic influenzas.
Airborne	Respiratory diseases acquired through transmission of pathogens through air (e.g., particles, droplets): for example, legionellosis. Includes respiratory infections that can be transmitted through air or other pathways, including infections transmitted through aerosols, fomites, or direct contact: Middle East respiratory syndrome coronavirus.

^aExamples are purposely not exhaustive and should be considered illustrative.

... COMO NÃO PODERIA DEIXAR DE SER!!!

Table 1. Determinants and drivers of infectious disease threat events, Europe, 2008–2013

Drivers, by group	Examples*
Globalization and environment	
Climate	Temperature, humidity, wind, rainfall. Can have an effect on exposure pathways of foodborne and waterborne diseases or the distribution of vectorborne diseases.
Natural environment	Land cover, vegetation, water ways, oceans, coastlines, water resources, land use, habitats, biodiversity. Can shift the distribution range and influence abundance of vectors (e.g., rodents, mosquitoes, ticks) as well as of host and reservoir animals.
Human-made environment	Urbanization, built environment, infrastructure, industries, intensive agriculture. Can enable propagation and dissemination of pathogens.
Travel and tourism	Movement of populations by automobile, train, ship, airplane. Can enable the importation of vectors, pathogens and infected persons into Europe and their dispersion within Europe.
Migration	Immigrant, emigrant, asylum seeker, settler. Can be vulnerable to or contribute to spread of infectious diseases in origin country, in transit, or in destination country.
Global trade	Import and export of goods and services across international boundaries via ship, airplane, rail, truck. Can result in the exportation or importation (on purpose or involuntarily) of host animals, disease vectors, or pathogens.
Sociodemographic	
Demographic	Population composition with regards to age, income, education. Can be associated with greater health vulnerabilities.
Social inequality	Uneven distribution of resources in society, including income, wealth, rights, privileges, social power, education. Disadvantaged groups can suffer disproportionately from infectious diseases.
Vulnerable groups	Children, premature infants, pregnant women, elderly persons, men who have sex with men, immunocompromised persons. Vulnerability can increase exposure and susceptibility to infectious diseases or decrease access to care and recovery.
Prevention	Childhood vaccination programs, adherence to treatment regimes, appropriate prescription practices. Distrust in prevention efforts can undermine control efforts (e.g., childhood vaccination programs. Neglect of prevention when traveling)
Lifestyle	High-risk behavior, such as intravenous drug use or unprotected sex with multiple partners. Can increase exposure and infection rates.
Occupational	Healthcare workers, veterinary and animal care personnel, butchers, farmers, cleaners. Lapses in infection control practices can put healthcare workers at risk.
Terrorism	Intentional release or dissemination of biologic agents. Intentional contamination of drinking water can result in community outbreaks.
Public health systems	
Healthcare system	European healthcare structure for the delivery of health services, including general practitioners, hospitals, clinics. Access to care, medicines, diagnostics, insurance coverage, for example, can affect health outcomes. Healthcare systems contribute to nosocomial infections.
Animal Health	Veterinary services, animal health and welfare measures, intensive livestock practices. High animal densities can promote infectious disease transmission. Infected animals close to human settlements can increase the risk for zoonotic epidemics.
Food and water quality	Agriculture, husbandry, farming, processing, handling, preparation and storage of food, man-made water systems (e.g., cooling towers, hot and cold water systems, spa pools, humidifiers), water treatment and distribution systems. Contamination of drinking and irrigation water sources and water distribution systems can result in both localized and community outbreaks.
Surveillance and reporting failure	Contamination of foodstuff along the chain from farm to fork can result in multistate epidemics. Systematic ongoing collection, collation, analysis, and dissemination of infectious disease data. Lapses in surveillance can impede a rapid response to infectious disease outbreaks. In contrast, increased surveillance will contribute to increased awareness and thus result in increased reporting of cases

*Examples are purposely not exhaustive and should be considered illustrative.

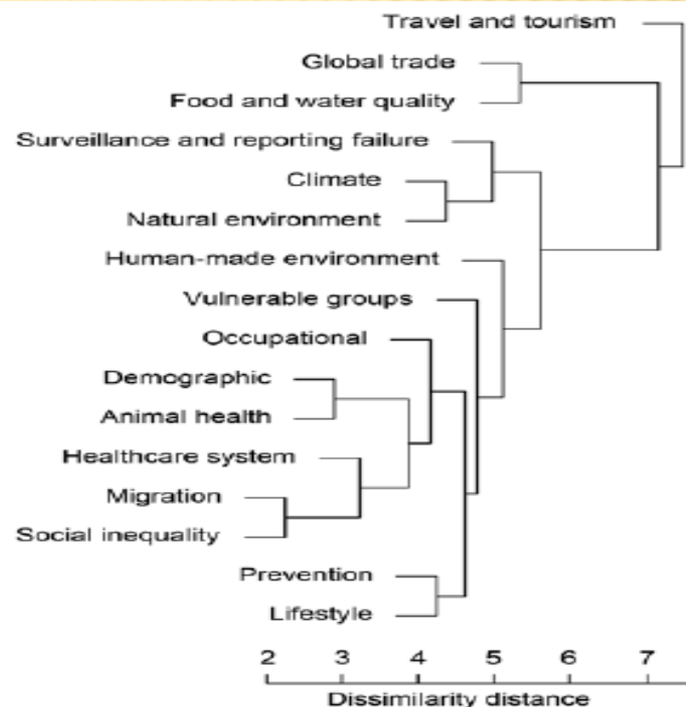


Figure 3. Cluster dendrogram from hierarchical cluster analysis of drivers contributing to observed infectious disease threat events (IDTEs), Europe, 2008–2013. Individual segments (leaves) on the lower part of the tree are more related to each other, as indicated by distances between the branches. Drivers below travel and tourism also occurred less often as underlying drivers of IDTEs and tended to be more contextual in nature. Scale bar indicates dissimilarity distance for drivers, as measured by frequency of pairwise co-occurrence in clusters. Similar drivers (e.g., that co-occurred in outbreaks) are at a close distance, and those that were more independent of other drivers show higher dissimilarity.

A IMPORTÂNCIA DO MEIO AMBIENTE...

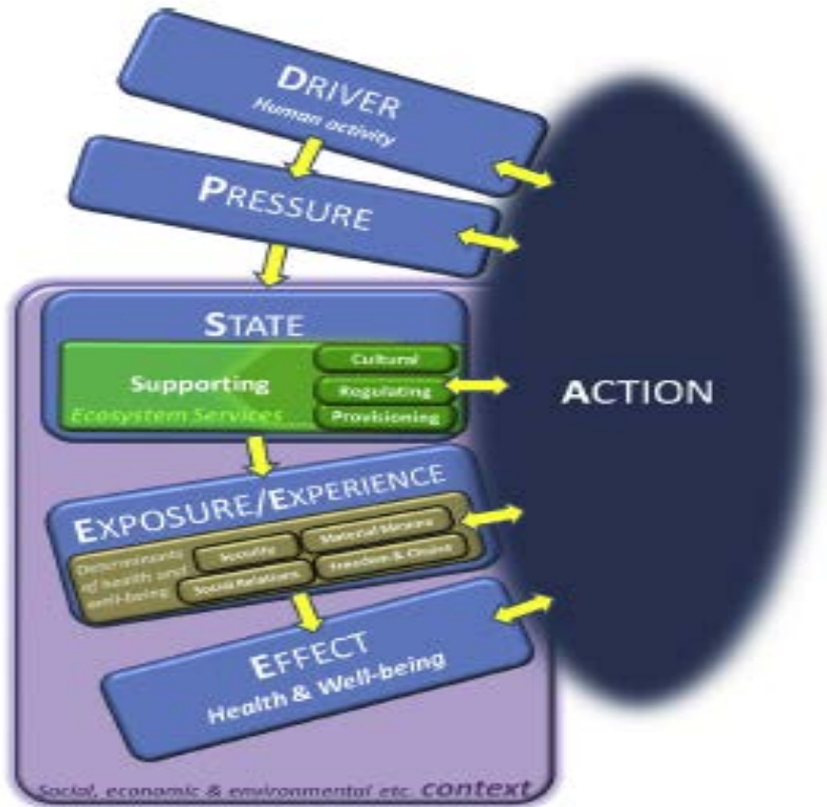


Fig. 1 – Ecosystem-enriched DPSEEA (eDPSEEA) – a conceptual framework for an integrated assessment of human and ecosystem health and ecosystem service provision.

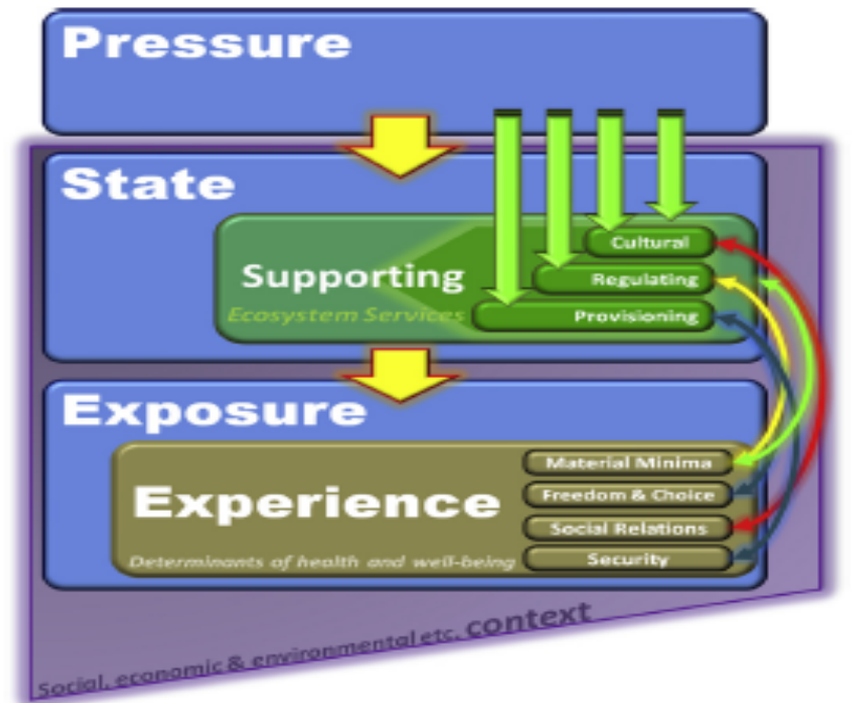


Fig. 2 – Illustrating the potential for feedback loops between Pressure, State and Exposure/Experience which is manifest when considering relationships between ecosystem services and determinants of human health and well-being. Feedbacks are depicted by two-directional arrows, but it should be noted that both positive and negative feedback effects may occur between a wide range of components of the eDPSEEA model.

... NOS DIVERSOS PROBLEMAS DE SAÚDE ...

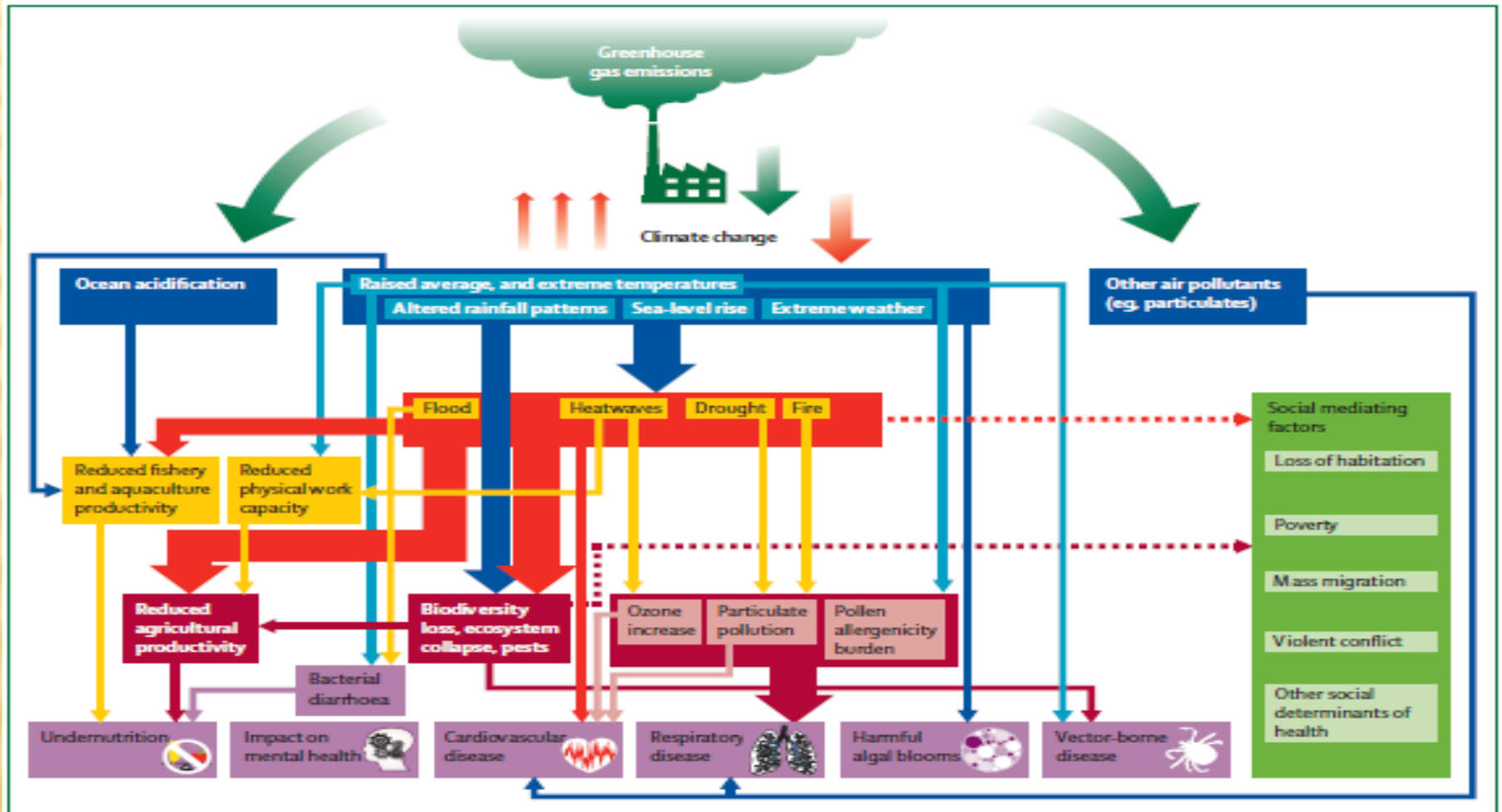


Figure 1: An overview of the links between greenhouse gas emissions, climate change, and health. The causal links are explained in greater detail in the section about climate change and exposure to health risks.

... ATRAVÉS DE EFEITOS DIRETOS E INDIRETOS ...

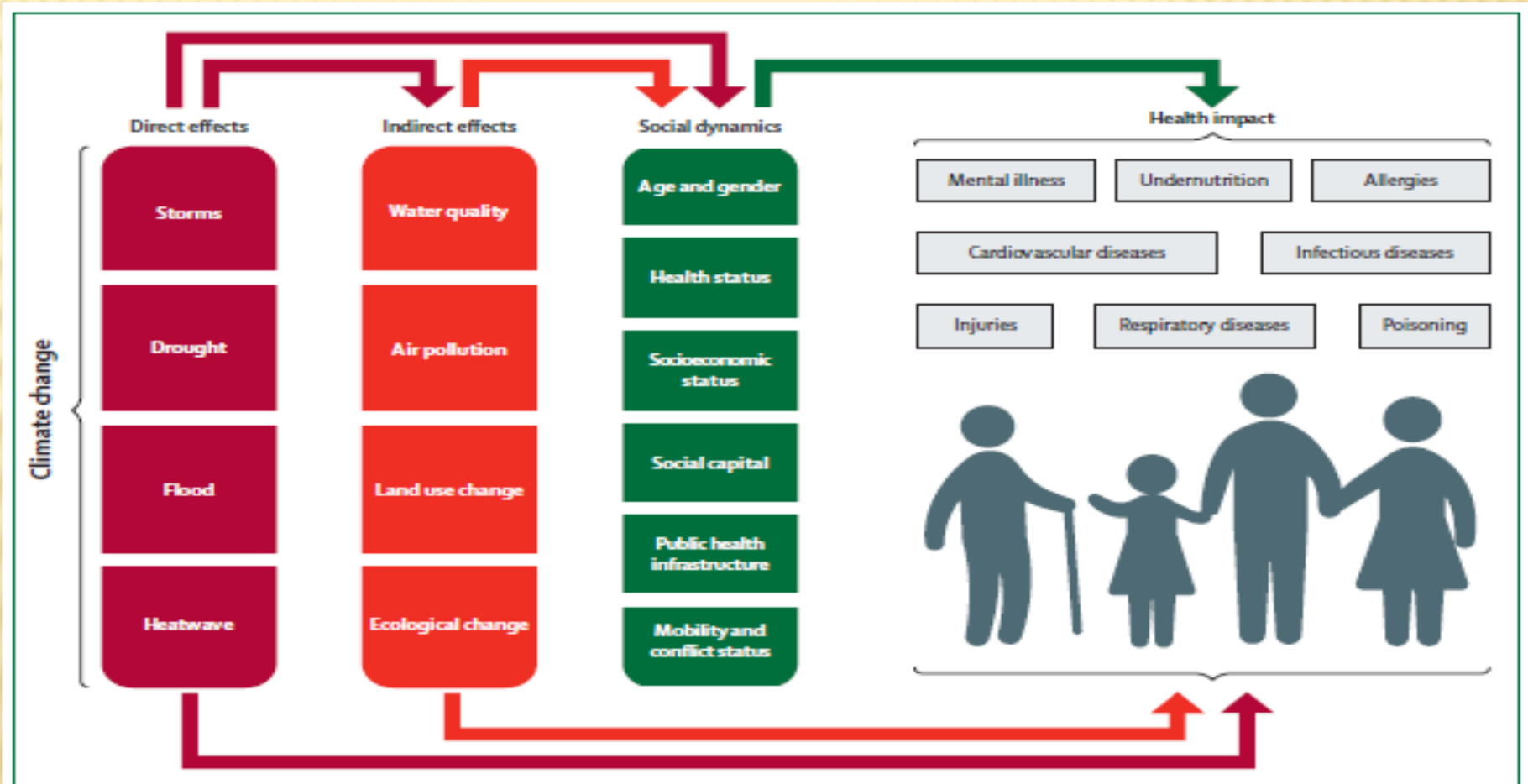


Figure 2: The direct and indirect effects of climate change on health and wellbeing

There are complex interactions between both causes and effects. Ecological processes, such as impacts on biodiversity and changes in disease vectors, and social dynamics, can amplify these risks. Social responses also ameliorate some risks through adaptive actions.

... QUE ABRANGE TODOS OS PAÍSES!!!

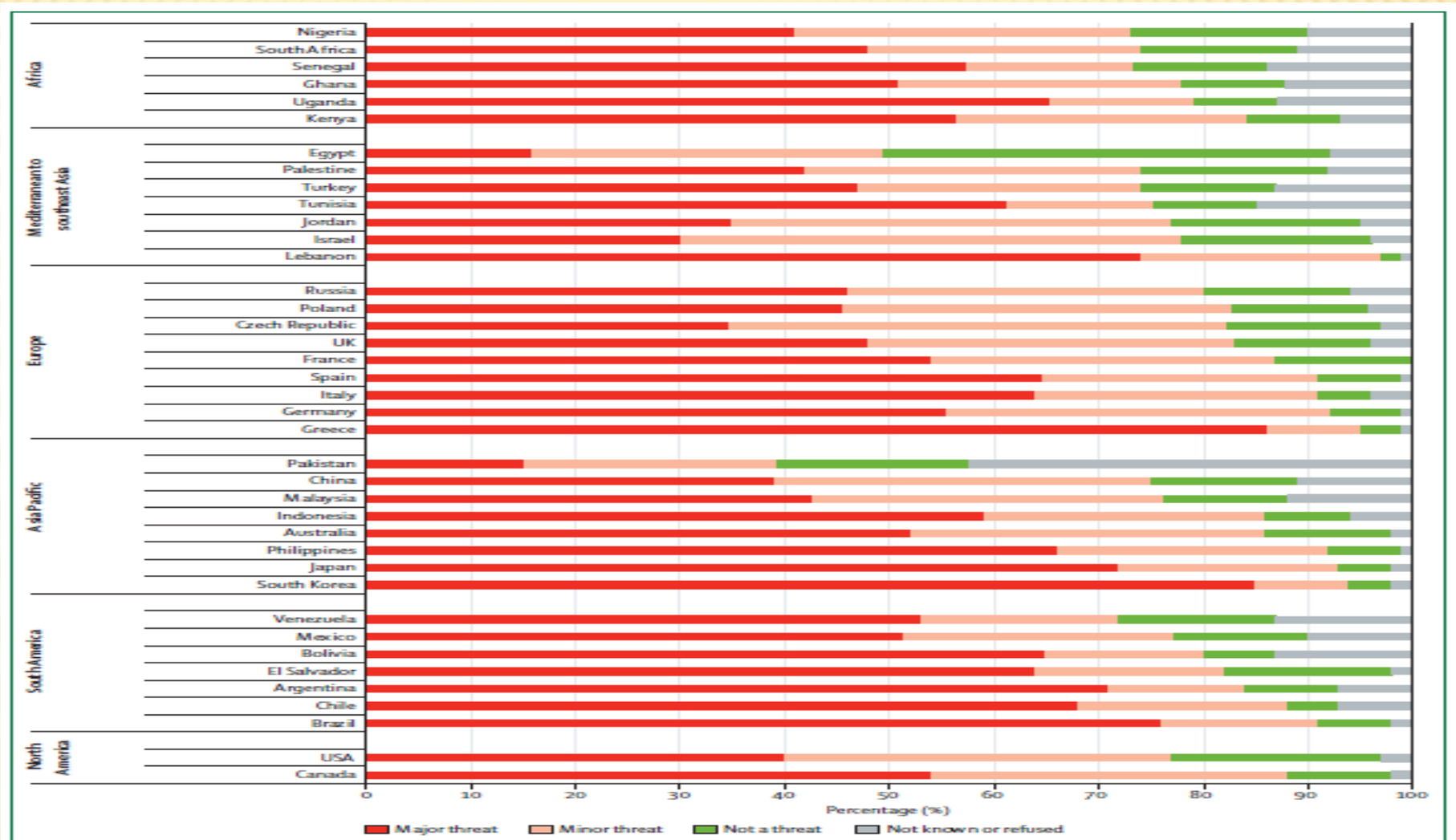


Figure 18: Perceptions of the threat of climate change, 2013**

IV)- O VIH/SIDA: OS DESAFIOS DA NOVA ESTRATÉGIA 90/90/90% DA ONUSIDA

A REALIDADE NACIONAL

FIGURA 3 ANOS POTENCIAIS DE VIDA PERDIDOS POR CAUSAS DE MORTE SELECIONADAS, EM PORTUGAL (2013)

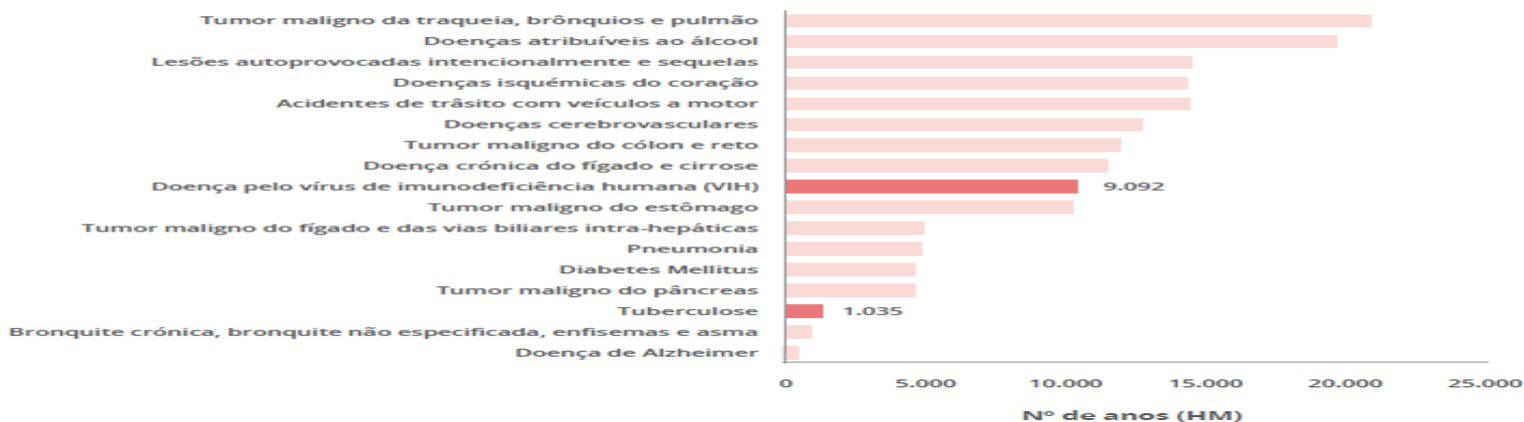
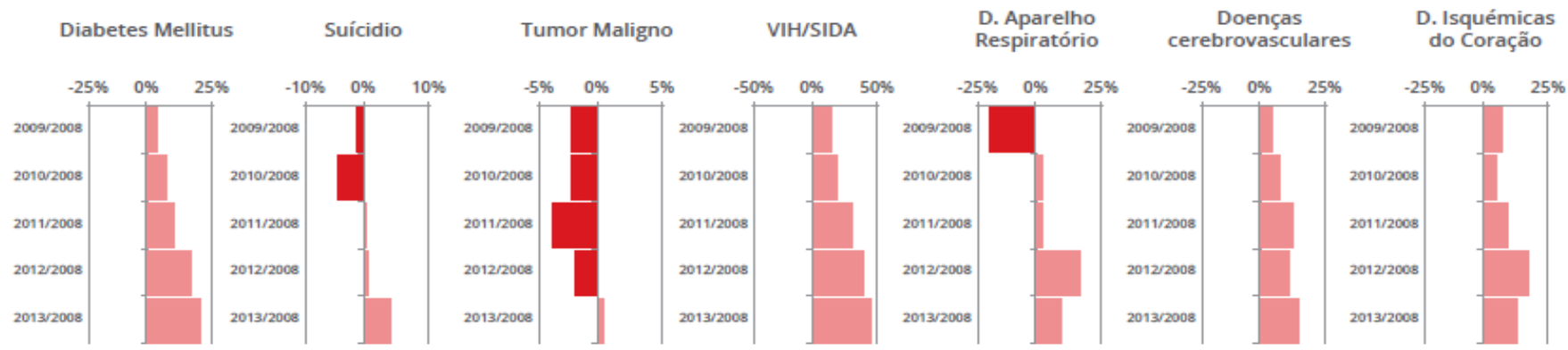
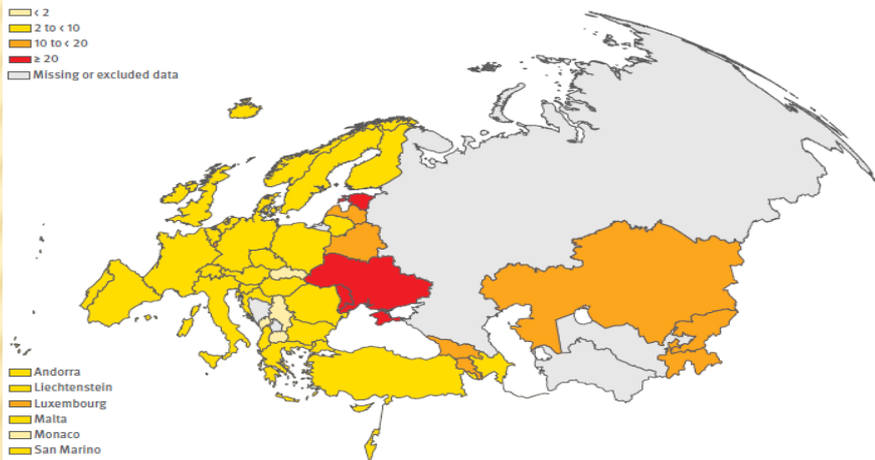


FIGURA 4 ANOS DE VIDA GANHOS, EM PORTUGAL, NO PERÍODO 2008-2013



O CASO PORTUGUÊS INSERIDO NO CONTEXTO EUROPEU

Map 1: New HIV diagnoses per 100 000 population, 2014



Map 7: Percentage of adult (>14 years) HIV diagnoses with CD4 <350 cells/mm³ at diagnosis, 2014

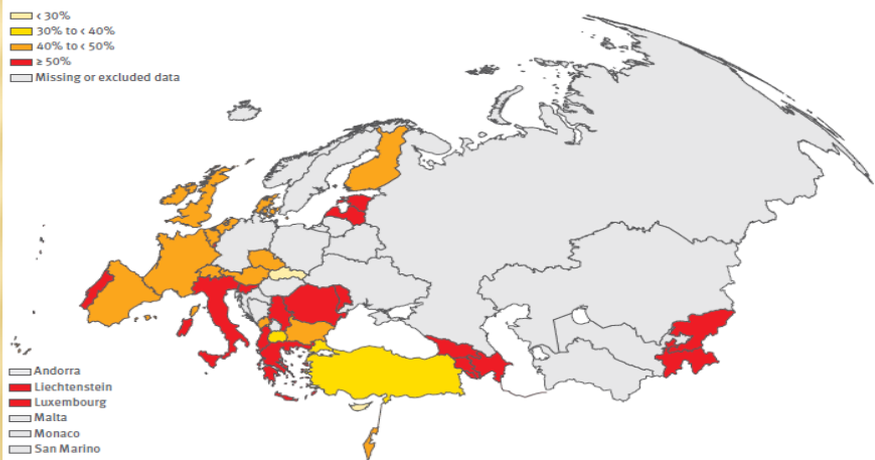
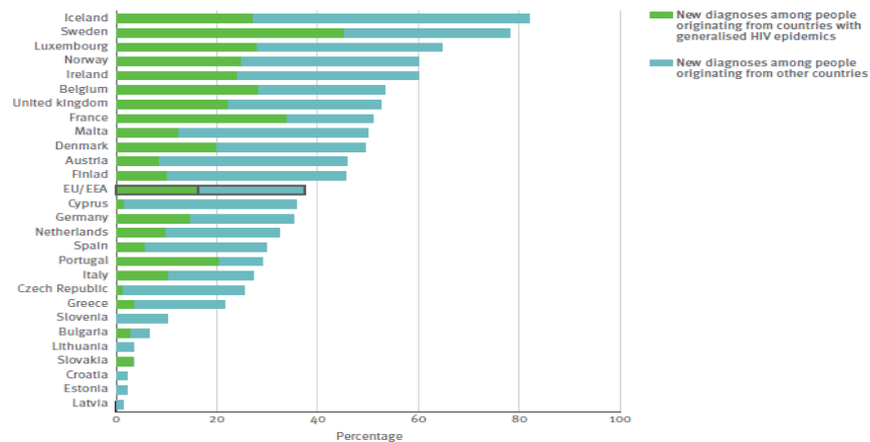
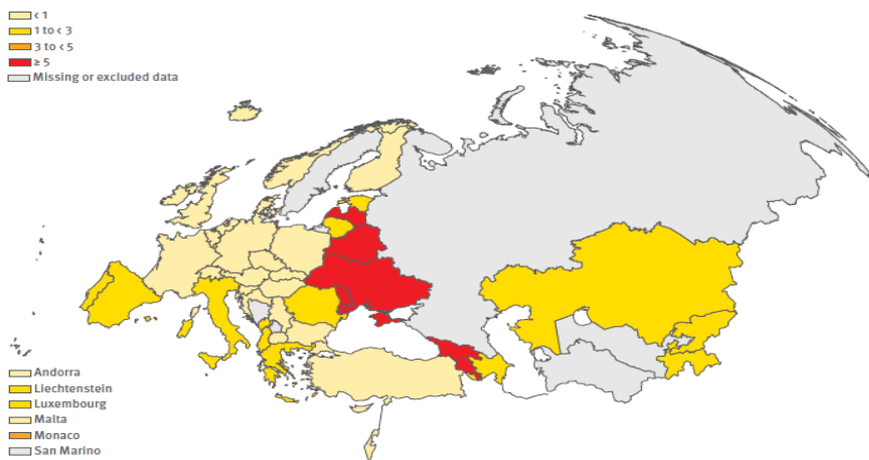


Figure 1.6: Percentage of new HIV diagnoses among migrants out of all reported cases with known information on region of origin, by country of report, EU/EEA, 2014 (n=25 525)



Map 8: AIDS diagnoses reported per 100 000 population, 2014



OS ENORMES IMPACTOS DA TARVC SOBRE A MORTALIDADE E A TRANSMISSÃO VERTICAL ...

BOSTON HEALTHCARE

White Paper | December 2012

Recognizing the Value of Innovation in HIV/AIDS Therapy

FIGURE 1. Evolution of the Treatment of HIV Infection: 1980s to Present

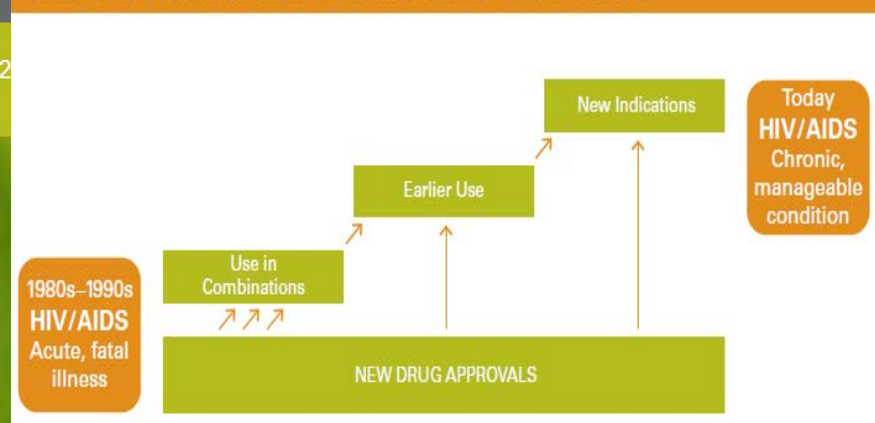


FIGURE 2. HIV Death Rates and HAART Treatment Advances (U.S., 1990-2010)

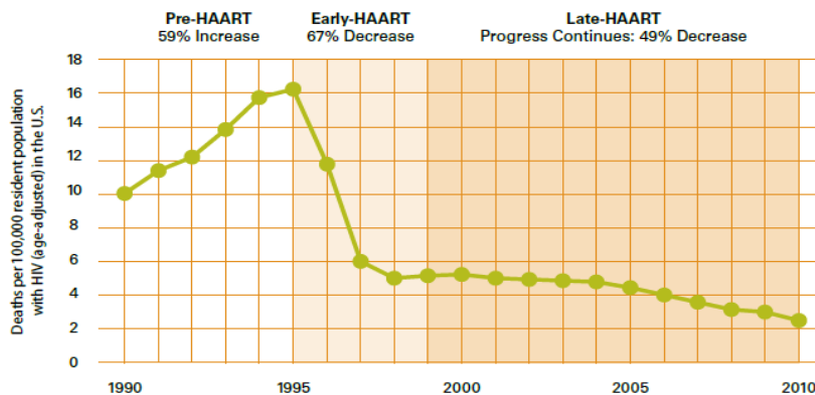
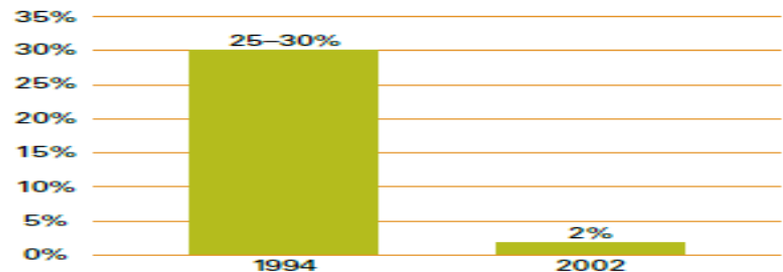


FIGURE 3. Reductions in Mother-to-Child HIV Transmission



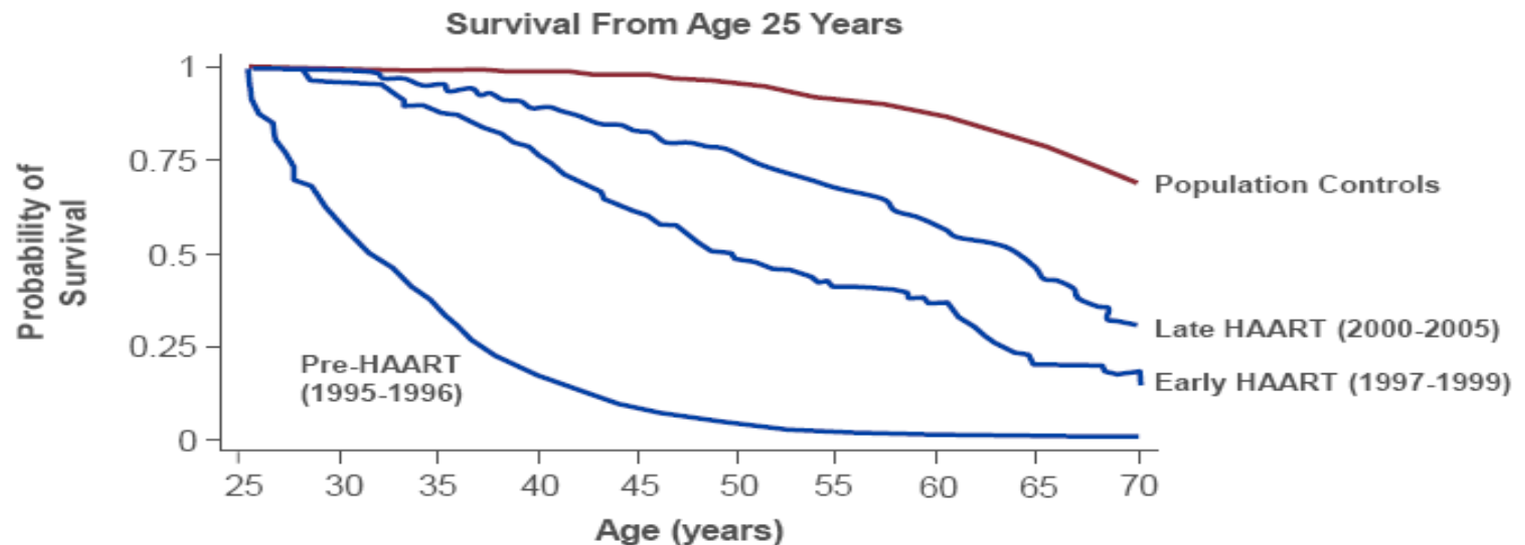
Source: Centers for Disease Control and Prevention, "Achievements in Public Health: Reduction in Perinatal Transmission of HIV Infection - United States, 1985-2005," MMWR, 55 (2 June 2006) 21, 592-597 <http://www.cdc.gov/mmwr/preview/mmwrhtml/mm5521a3.htm>.

... E SOBRE A SOBREVIVÊNCIA DOS DOENTES

Survival trends in HIV infection have changed since the adoption of cART

UP TO THE CHALLENGE
OF LONG-TERM TREATMENT SUCCESS

Cumulative survival curve for HIV-infected persons (non-HCV co-infected) and persons from the general population.



n=383,862 (HIV-infected patients, n=3,990;
General population controls, n=379,872)

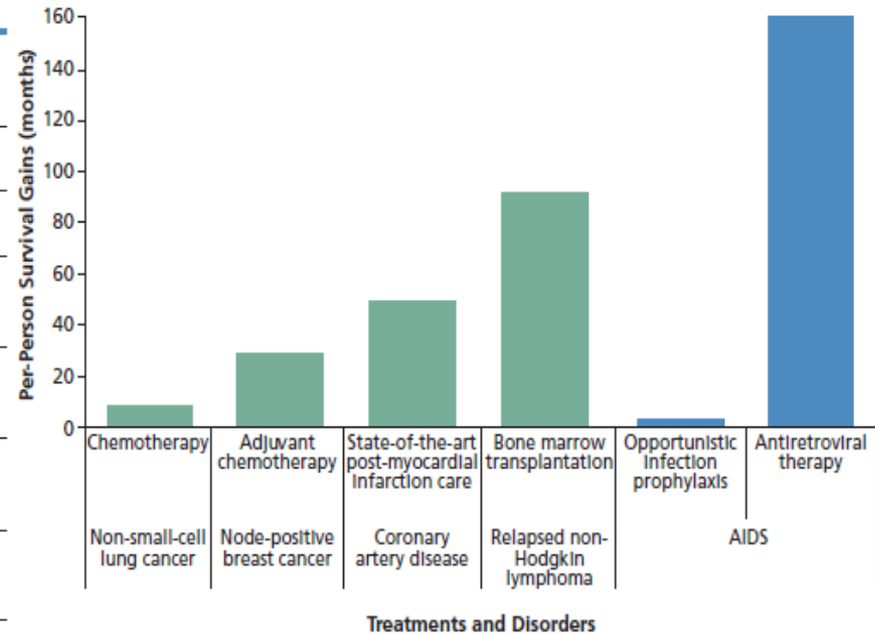
A SUA DEMONSTRADA CUSTO-EFETIVIDADE ...

Perspective

Cost-Effectiveness of HIV Interventions: From Cohort Studies and Clinical Trials to Policy

Table 1. Per-Person Survival Benefit in AIDS Patients by Treatment Era

Years	Intervention	Per-Person Survival Benefit (months)	No. of Patients Diagnosed and Entering Care	Total Survival Benefit (years)
1989-1992	PCP prophylaxis	3.1	158,370	40,912
1993-1995	PCP prophylaxis + MAC prophylaxis	24.4	226,458	460,465
1996-1997	PCP prophylaxis + MAC prophylaxis + ART Era 1	93.7	72,716	567,788
1998-1999	PCP prophylaxis + MAC prophylaxis + ART Era 2	132.6	52,702	582,359
2000-2002	PCP prophylaxis + MAC prophylaxis + ART Era 3	138.8	71,946	832,179
2003	PCP prophylaxis + MAC prophylaxis + ART Era 4	159.9	24,780	330,189
Total				2,813,892



ART indicates antiretroviral therapy; Eras 1-4, periods characterized by improvements in antiretroviral therapy over time; MAC, *Mycobacterium avium* complex; PCP, *Pneumocystis jirovecii* pneumonia. Adapted from Walensky et al, *J Infect Dis*, 2006.

Figure 1. Per-person survival gains with treatment in patients with AIDS compared with gains associated with interventions for other common diseases in the United States. Adapted from Walensky et al, *J Infect Dis*, 2006.

... BEM COMO QUANTO AO DIAGNÓSTICO PRECOCE ...

Table 2. Cost-Effectiveness Ratios for Select HIV-Related Interventions and for HIV and Non-HIV-Related Screening Interventions

Intervention	Drug	Cost-Effectiveness Ratio (\$/QALY)*	Reference
HIV Interventions			
PCP/Pneocystis prophylaxis	TMP-SMX	\$2600	Freedberg et al, JAMA, 1998
Antiretroviral therapy	Zidovudine/ lamivudine/ efavirenz	\$13,000	Freedberg et al, N Engl J Med, 2001
Genotypic resistance test, at treatment failure	NA	\$17,900	Weinstein et al, Ann Intern Med, 2001
Genotypic resistance test, treatment-naive	NA	\$20,200	Sax et al, Clin Infect Dis, 2005
Inpatient HIV screening	NA	\$15,100	Walensky et al, Am J Med, 2005
MAC prophylaxis	Azithromycin	\$44,500	Freedberg et al, JAMA, 1998
HIV and Other Screening Interventions			
HIV screening every 5 years in patients at high risk	NA	\$42,200	Paltiel et al, N Engl J Med, 2005
Breast cancer screening: annual mammogram, 50-69 years old	NA	\$57,500	Salemann et al, Ann Intern Med, 1997
Colon cancer screening: FOBT + sigmoidoscopy every 5 years, 50-85 years old	NA	\$53,600	Frazier et al, JAMA, 2000
Type 2 diabetes: one-time FPG, > 25 years old	NA	\$63,000	CDC, JAMA, 1998

CDC indicates Centers for Disease Control and Prevention; FOBT, fecal occult blood test; FPG, fasting plasma glucose; MAC, Mycobacterium avium complex; NA, not applicable; PCP, Pneumocystis jirovecii pneumonia; TMP-SMX, trimethoprim-sulfamethoxazole. *All costs adjusted to 2001 US dollars. Ratios from Walensky et al, Am J Med, 2005.

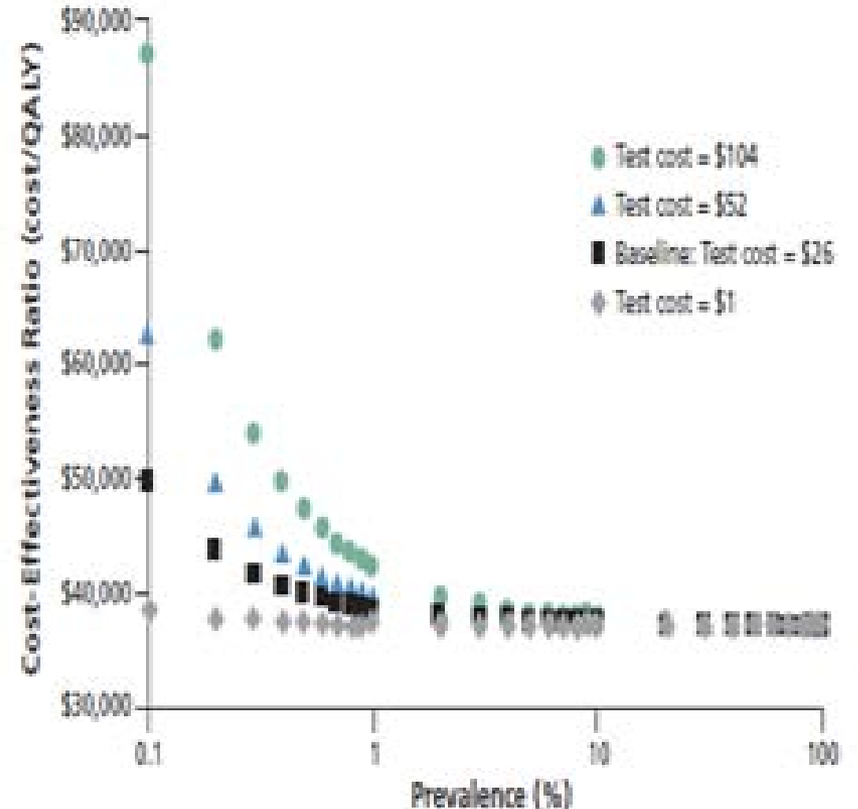
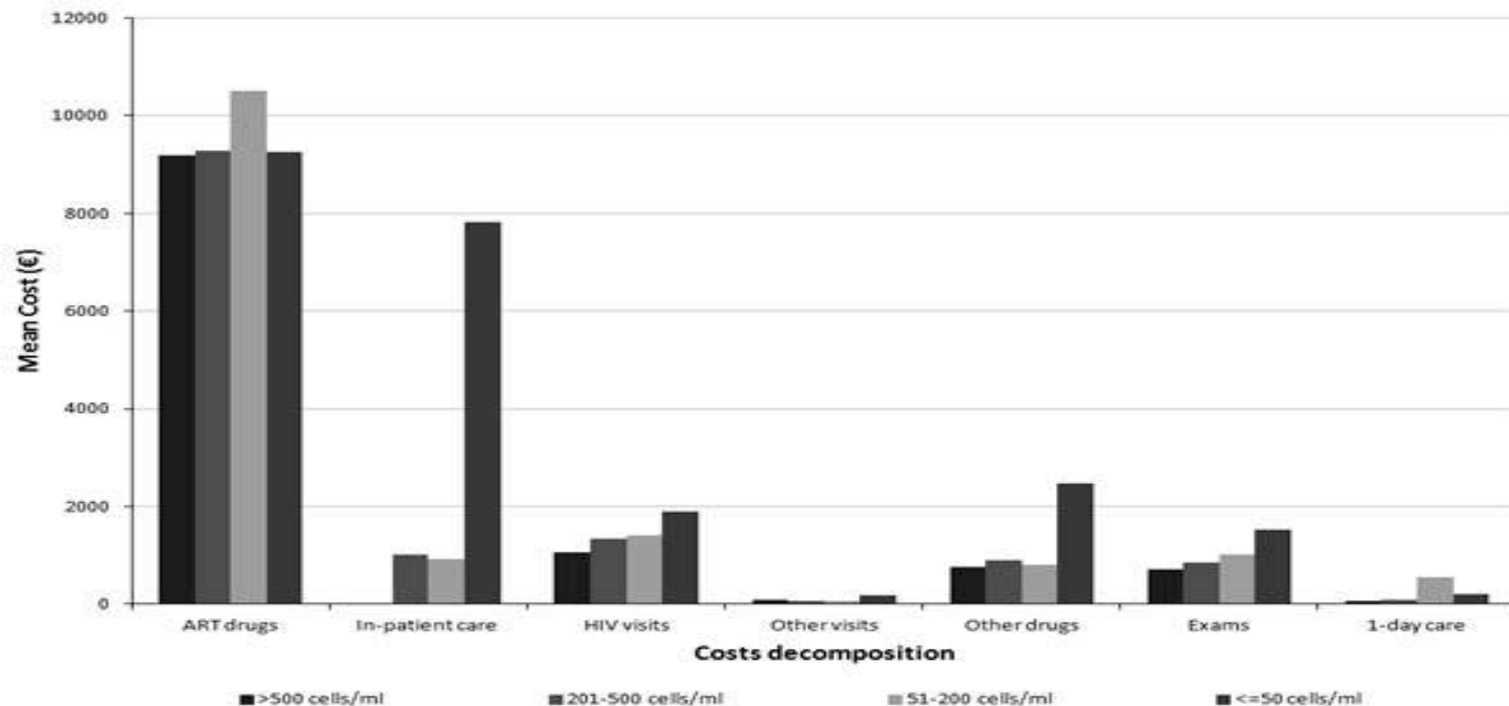


Figure 2. Cost-effectiveness ratios of HIV screening according to prevalence of undiagnosed HIV infection in the testing population and different hypothetical test costs. QALY indicates quality-adjusted life-years. Adapted from Walensky et al, Am J Med, 2005.

... PREMISSE QUE TAMBÉM JÁ FOI DEMONSTRADA EM PORTUGAL!!!

DIRECT TREATMENT COSTS OF HIV/AIDS IN PORTUGAL

Julian Perelman^{1,2}, Joana Alves¹, Ana Cláudia Miranda³, Céu Mateus^{1,2}, Kamal Mansinho^{2,3}, Francisco Antunes⁴, Joaquim Oliveira⁵, José Poças⁶, Manuela Doroana⁴, Rui Marques⁷, Eugénio Teófilo⁸, João Pereira^{1,2}



0 CASO DOS EUA ...

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PLOS MEDICINE

Review

HIV Treatment as Prevention: The Utility and Limitations of Ecological Observation

M. Kumi Smith¹, Kimberly A. Powers^{1,2}, Kathryn E. Muessig², William C. Miller^{1,2}, Myron S. Cohen^{1,2,3*}

¹ Department of Epidemiology, University of North Carolina, Chapel Hill, North Carolina, United States of America, ² Department of Medicine, University of North Carolina, Chapel Hill, North Carolina, United States of America, ³ Department of Microbiology and Immunology, University of North Carolina, Chapel Hill, North Carolina, United States of America

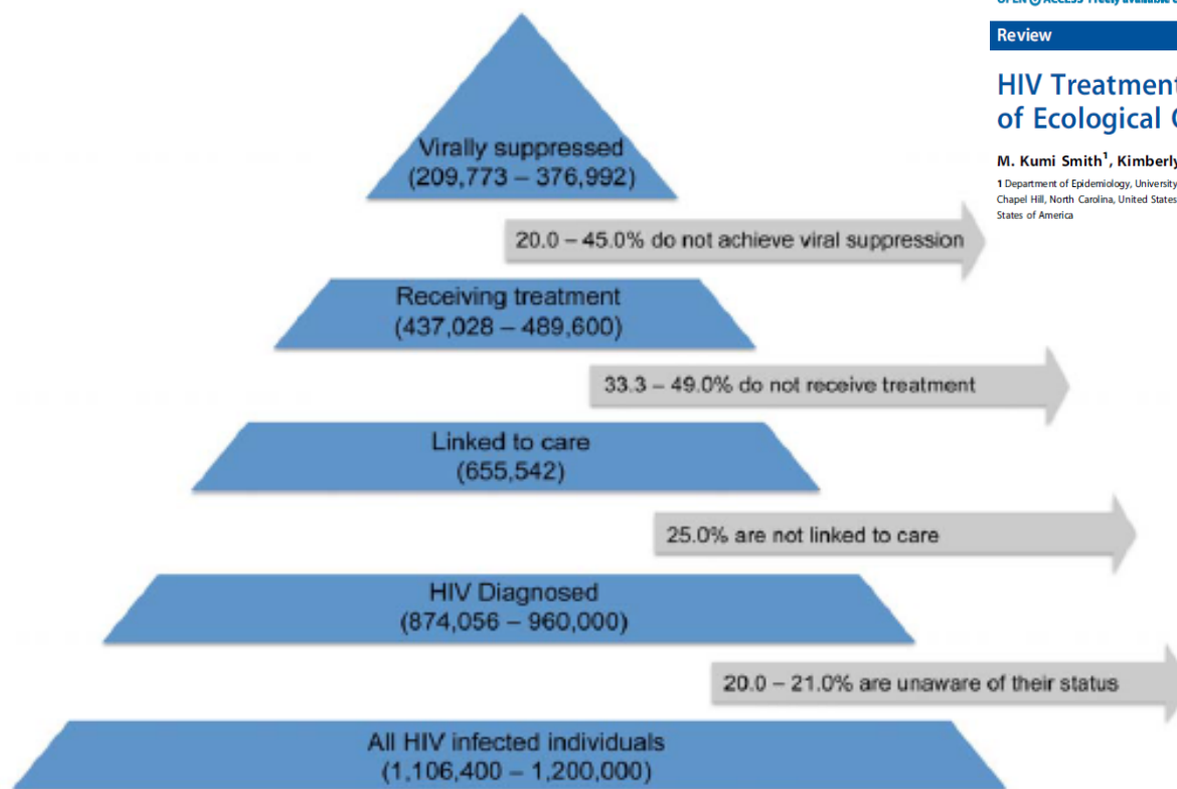


Figure 1. Estimated numbers of HIV-infected individuals in the US retained (and corresponding percentages lost) at various stages of the test, link, and treat cascade. This figure is based on data from [61,62].

doi:10.1371/journal.pmed.1001260.g001

OS ESTUDOS FUNDAMENTAIS E AS NOVAS RECOMENDAÇÕES TERAPÊUTICAS

The NEW ENGLAND
JOURNAL of MEDICINE

ESTABLISHED IN 1812 AUGUST 27, 2015 VOL. 373 NO. 9

Initiation of Antiretroviral Therapy in Early Asymptomatic HIV Infection

The INSIGHT START Study Group*



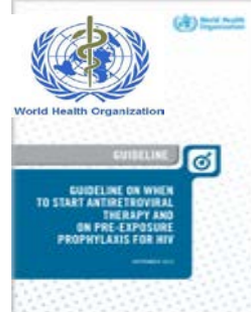
The NEW ENGLAND
JOURNAL of MEDICINE

ESTABLISHED IN 1812 NOVEMBER 30, 2006 VOL. 355 NO. 22

CD4+ Count-Guided Interruption of Antiretroviral Treatment

The Strategies for Management of Antiretroviral Therapy (SMART) Study Group*

BILL & MELINDA GATES foundation



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PLOS MEDICINE

Review

HIV Treatment as Prevention: Models, Data, and Questions—Towards Evidence-Based Decision-Making

The HIV Modelling Consortium Treatment as Prevention Editorial Writing Group*

Health benefits, costs, and cost-effectiveness of earlier eligibility for adult antiretroviral therapy and expanded treatment coverage: a combined analysis of 12 mathematical models

Jeffrey W Eaton*, Nicolas A Menzies*, John Stover, Valentina Cambiano, Leonid Chindelevitch, Anne Cori, Jan A C Hontela, Salal Humair, Cliff C Kerr, Daniel J Klein, Sharmistha Mishra, Kate M Mitchell, Brooke E Nichols, Peter Vickerman, Roel Bakker, Till Barnighausen, Anna Bershteyn, David E Bloom, Marie-Claude Bolly, Stewart T Chang, Ted Cohen, Peter J Dodd, Christophe Fraser, Chaitra Gopalappa, Jens Lundgren, Natasha K Martin, Evdinn Mikkelsen, Elisa Mountain, Quang D Pham, Michael Pickles, Andrew Phillips, Lucy Platt, Carel Pretorius, Holly J Prudden, Joshua A Salomon, David A M C van de Vijver, Sake J de Vlas, Bradley G Wagner, Richard G White, David P Wilson, Lei Zhang, John Blandford, Gesine Meyer-Rath, Michelle Remme, Paul Revill, Nalinee Sangrujee, Fern Terris-Prestholt, Meg Doherty, Nathan Shaffer, Philippa Eastbrook, Gottfried Hirschall, Timotij B Hallett

A IMPORTÂNCIA DA IMPLEMENTAÇÃO DE UMA ESTRATÉGIA COMBINADA



OPEN ACCESS Freely available online
Review

PLOS MEDICINE

HIV Treatment as Prevention: Considerations in the Design, Conduct, and Analysis of Cluster Randomized Controlled Trials of Combination HIV Prevention

Marie-Claude Boily^{1*}, Benoit Masse², Ramzi Alsallaq³, Nancy S. Padian^{4,5}, Jeffrey W. Eaton¹, Juan F. Vesga¹, Timothy B. Hallett¹

¹Department of Infectious Disease Epidemiology, School of Public Health, Imperial College London, London, United Kingdom, ²CHU Sainte-Justine Research Centre, University of Montreal, Montreal, Quebec, Canada, ³College of Nursing Global, New York University, New York, United States of America, ⁴Office of the US Global AIDS Coordinator, US Department of State, Washington, District of Columbia, United States of America, ⁵Department of Epidemiology, University of California, Berkeley, California, United States of America

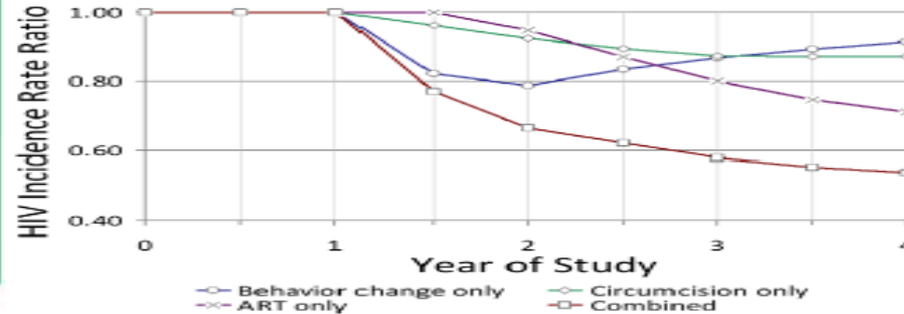
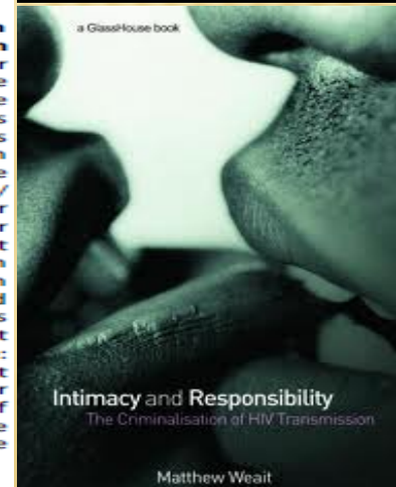
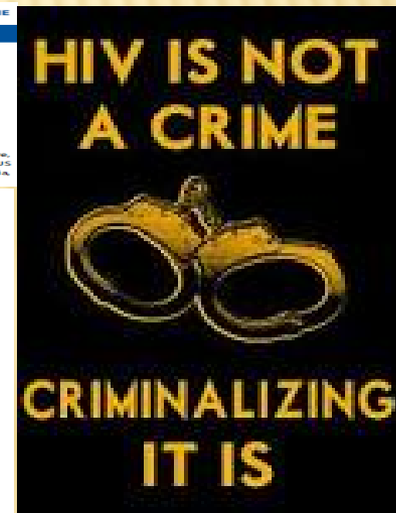


Figure 1. Predicted short-term impact of three intervention components linked to HIV testing in KwaZulu-Natal, South Africa. The model is based on a high-transmission setting under conditions of the current standard of care versus a high-coverage combination intervention (see [26]). The instantaneous HIV incidence rate ratio in the y-axis is intervention versus control. Impact estimates include an initial 6-mo period of preparation for the study. Assumptions for the combination intervention: 90% of adults in the intervention community are tested in the first year and thereafter every 4 y; those who test positive reduce risk behaviour for 3 y (on average) (25.0%/12.5% of men/women increase condom use; 25%/25% reduce partner acquisition); 70% of uncircumcised men are circumcised in the first year (efficacy = 60%); and all those in need of treatment (CD4 cell count <350 cells/ μ l) are immediately treated with ART (efficacy = 92%) with an annual dropout rate from treatment of 5%. The efficacy of MC in reducing susceptibility is assumed to be immediate (i.e., the wound healing period is negligible). Viral suppression for infected individuals once on treatment is immediate (i.e., no delay between treatment initiation and viral suppression). Assumptions for the standard of care: 20% of individuals test annually; 12.5%/6.5% of men/women who test positive increase condom use, and 12.5%/12.5% reduce partner acquisition, for one year; HIV-positive individuals are treated if CD4 <200 cells/ μ l (dropout rate of 15%); and 27% of men are circumcised at baseline and 10% more over 4 y since the start of the intervention.

doi:10.1371/journal.pmed.1001250.g001



FOI PENA QUE...



Ist Porto Meeting in
Mathematics and Biology

Project on Epidemiological
and Economical Aspects of
HIV/AIDS transmission in
Portugal

- ✗ Estudo onde se pretendia analisar:
 - + 1)- CV comunitária
 - + 2)- CV cumulativa individual



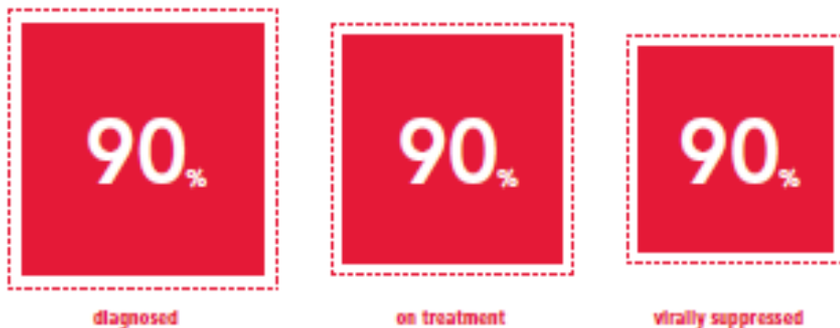
A ESTRATÉGIA 90%-90%-90% DA UNAIDS

90-90-90

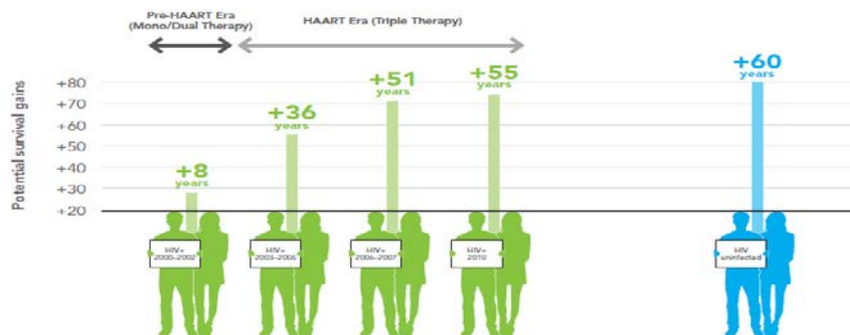


An ambitious treatment target to help end the AIDS epidemic

THE TREATMENT TARGET



HIV TREATMENT CAN NORMALIZE SURVIVAL



Expected impact of HIV treatment in survival of a 20 years old person living with HIV in a high income setting (different periods)

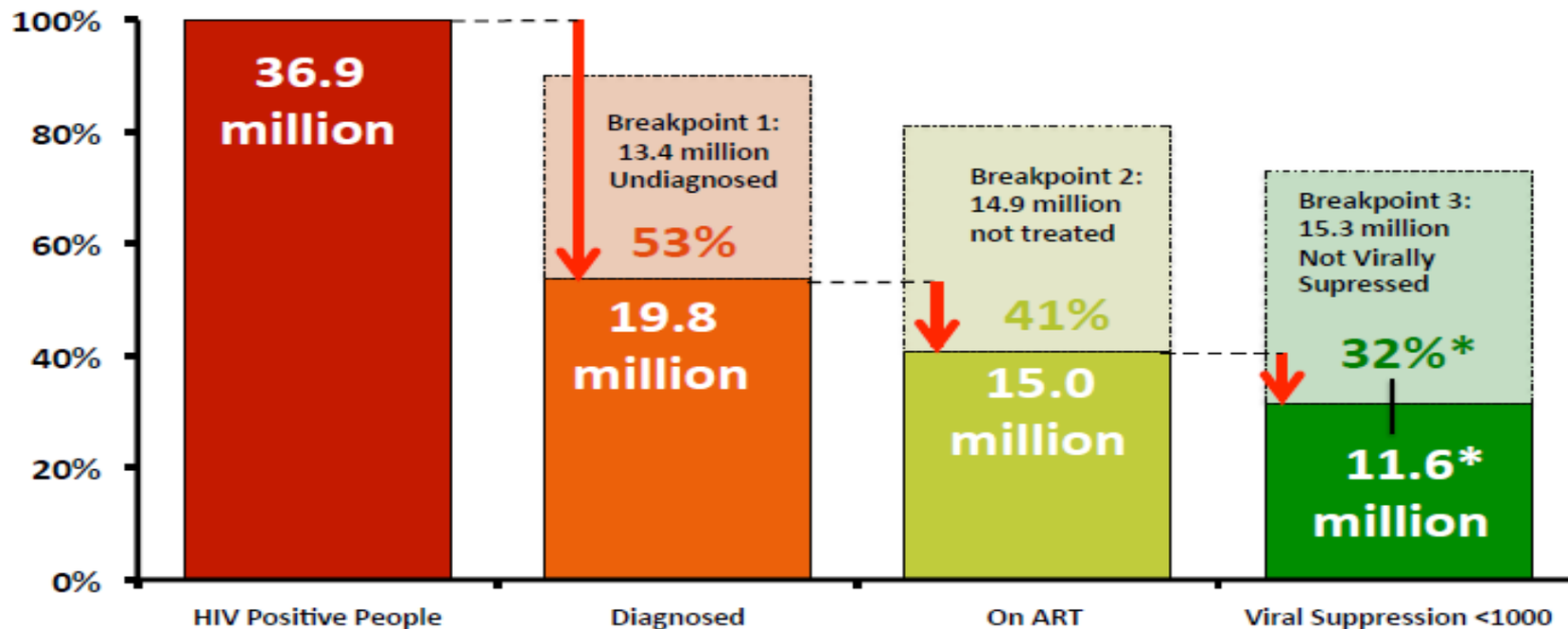
Source: Samji H et al., PLoS ONE, 2013.

THE CHOICES



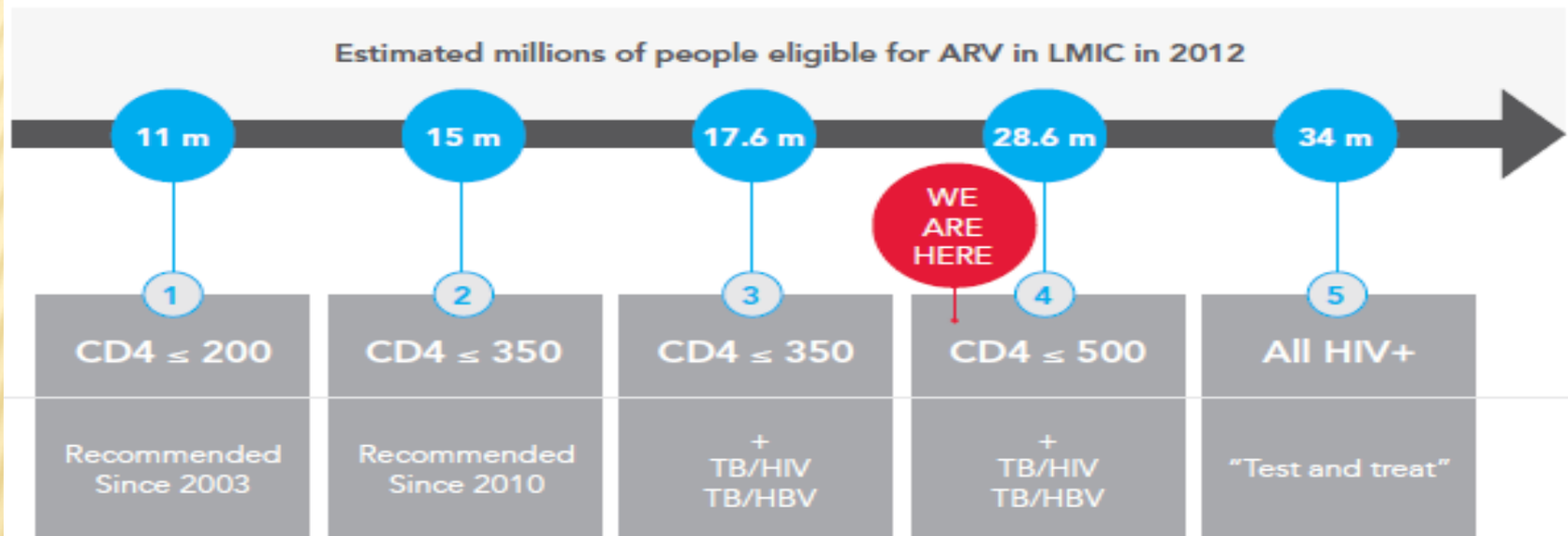
AS DIFERENÇAS ENTRE AS INTENÇÕES E A REALIDADE

Global Estimates (2014-15) vs the Gap to reach 90-90-90 Targets



A MAGNITUDE EFETIVA DESTA ESTRATÉGIA

SCENARIOS OF ANTIRETROVIRAL TREATMENT ELIGIBILITY: WHO VISION



WE ARE HERE

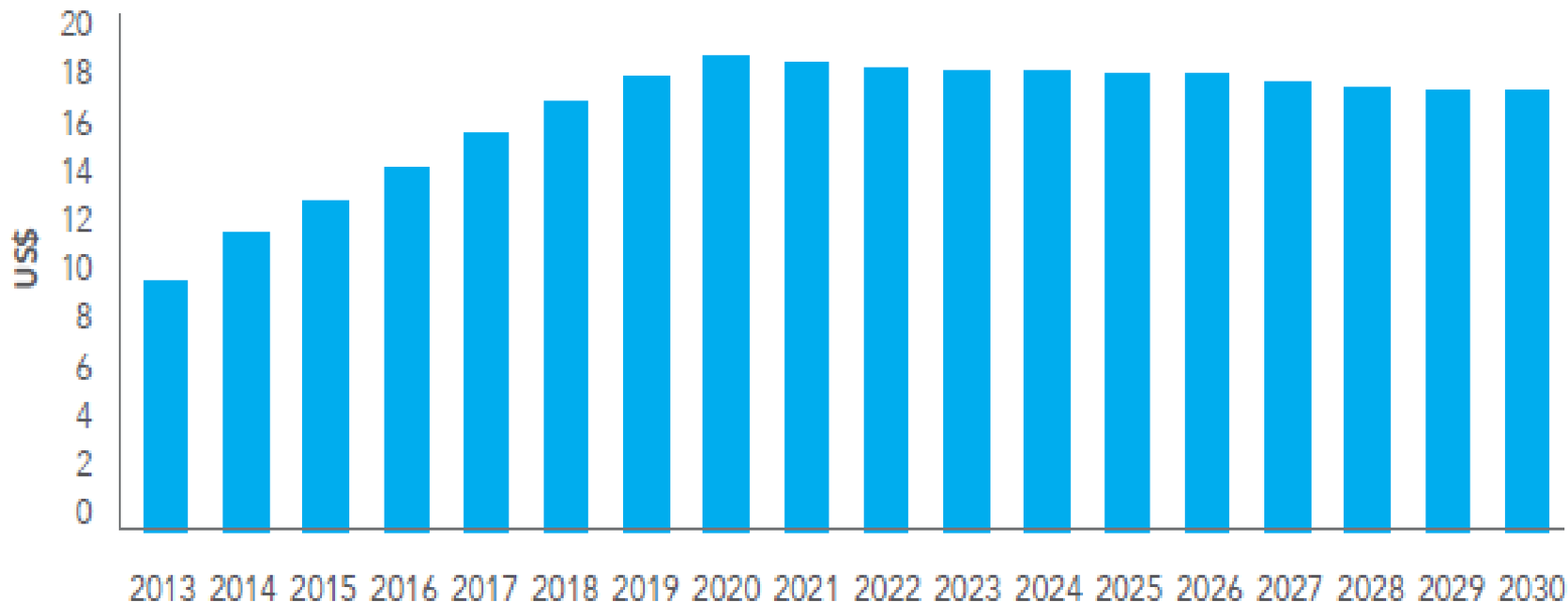
Scenarios of ARV eligibility

ART regardless of CD4 count for:

- Serodiscordant couples
- Pregnant women
- Children < 5 years

SENDO CERTO QUE OS CUSTOS DIRETOS SERÃO ENORMES, OS DA NÃO IMPLEMENTAÇÃO DESTA ESTRATÉGIA SERÃO, CONTUDO, A PRAZO, MUITO SUPERIORES

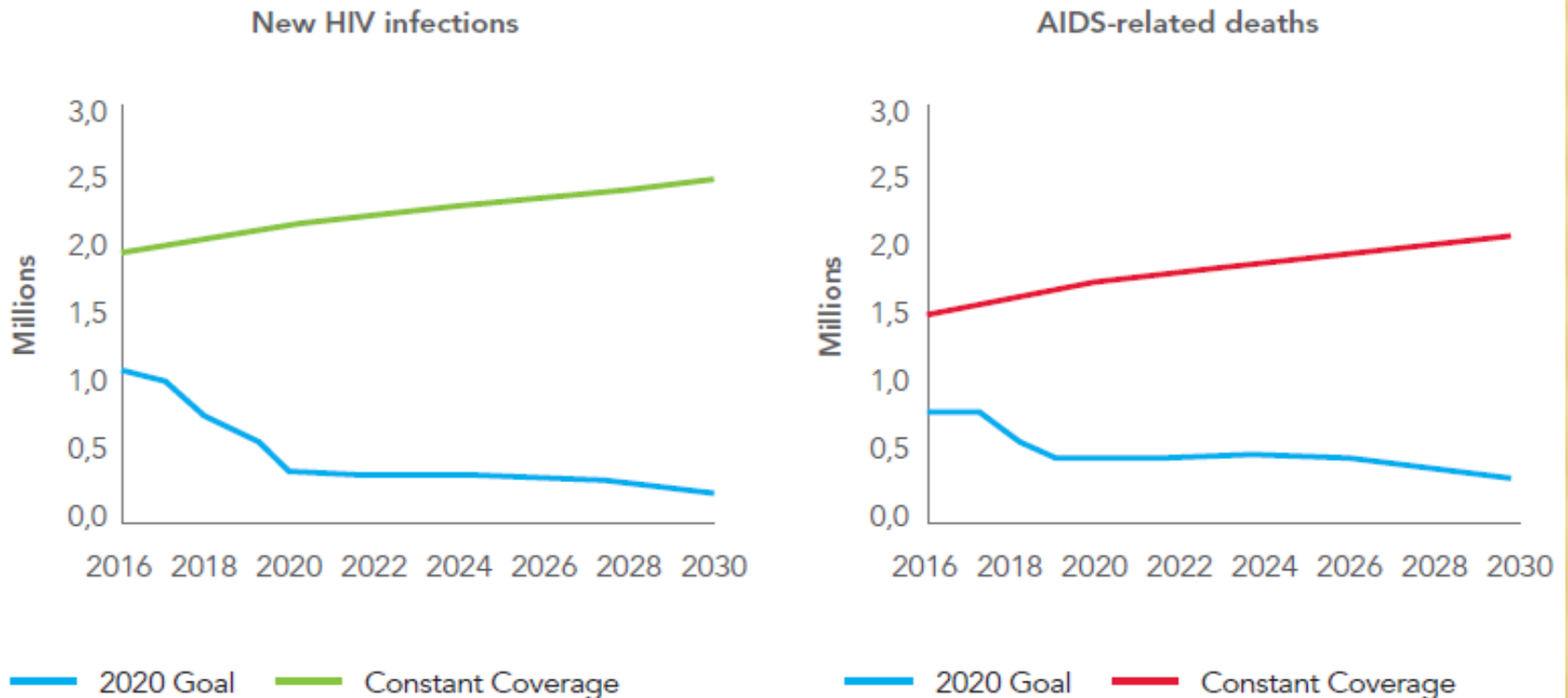
RESOURCE NEEDS FOR THE 90-90-90 TREATMENT TARGET, 2016-2030 (DRUGS, SERVICE DELIVERY, TESTING AND COUNSELLING, COMMUNITY MOBILIZATION AND PRE-ART COSTS)



Source: UNAIDS Global Price Tag Estimates, September 2014. Unpublished results.

OS OBJETIVOS E OS IMPACTOS

IMPACT OF THE 90-90-90 TARGET ON HIV INFECTIONS AND AIDS-RELATED DEATHS, 2016-2030



Source: The Gap Report, UNAIDS, 2014.

A HOMENAGEM DEVIDA: “O SEU A SEU DONO”!!!



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Published in final edited form as:

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Treatment as Prevention—Where Next?

Mark Hull^{1,2}, Joep Lange^{3,*}, and Julio S.G. Montaner^{1,2}



¹BC Centre for Excellence in HIV/AIDS, Vancouver, Canada ²University of British Columbia, Vancouver, Canada ³University of Amsterdam, the Netherlands

I REMOS SER CAPAZES?



IAS 2015
vancouver, canada
18th IAS Conference on HIV Pathogenesis,
Treatment & Prevention 19-22 July 2015
IAS2015.ORG



Code: MOAD01, MOAD0102
Title: 90-90-90: Delivering on the Targets
Date: Monday, 20 July 2015
Time: 16:30-18:00
Room: Ballroom C-D

Can the UNAIDS 90-90-90 target be reached?

Analysis of national HIV treatment cascades

Jacob Levi¹ & Alice Raymond¹; Anton Pozniak²; Pietro Vernazza³; Philipp Kohler³; Andrew Hill²

V)- A REALIDADE AO NÍVEL DO CHS HSB DE SETÚBAL

A PARTICIPAÇÃO NA REDE DE VIGILÂNCIA DE DOENÇAS TRANSMISSÍVEIS VETORIAIS DO CEVDI DO INSA ...

REVIVE 2011-2015 Culicídeos e Ixodídeos

Rede de Vigilância de Vetores

Instituto Nacional de Saúde Doutor Ricardo Jorge
Administrações Regionais de Saúde
Instituto de Administração de Saúde e Assuntos Sociais
Direção-Geral de Saúde

Centro de Estudos de Vetores e Doenças Infecciosas Doutor Francisco Cambournac

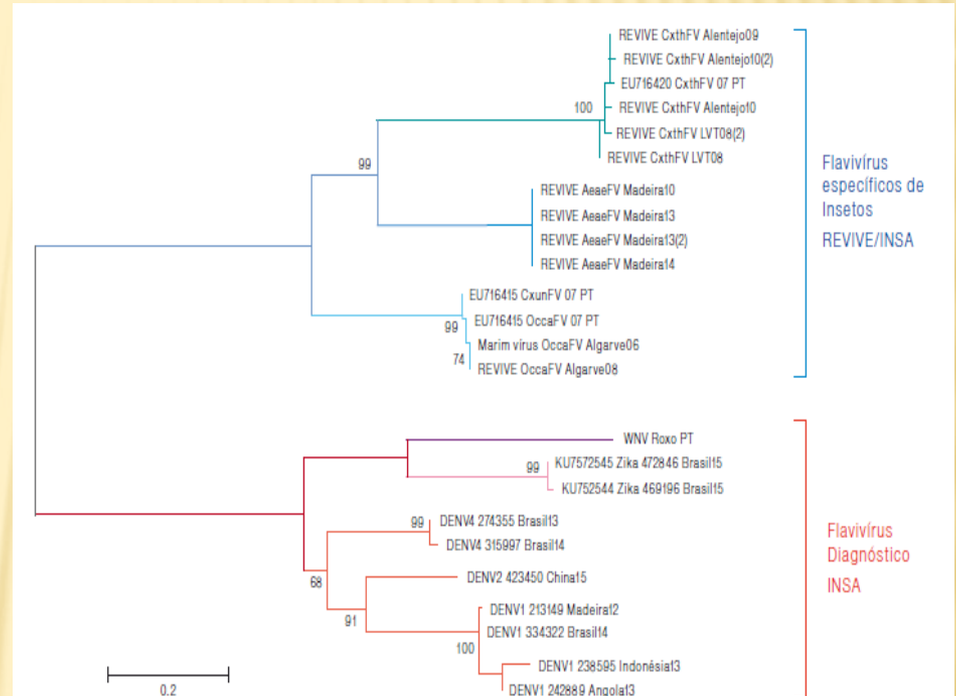


Figura 14: Árvore filogenética máxima parcimónia baseada em sequências parciais da proteína não estrutural NS5 obtida recorrendo ao software *Molecular Evolutionary Genetics Analysis* (MEGA), versão 6. Os valores de bootstrap superiores a 60 são apresentados na árvore. Os flavivírus específicos de insetos estão representados a azuis e os flavivírus patogénicos para o Homem identificados no INSA (no âmbito do diagnóstico laboratorial) estão representados a vermelhos. Todas as sequências apresentam a identificação do vírus, país ou região de origem e ano de deteção: AaaaFV, Flavivírus detetados em *Aedes aegypti*; CxthFV, Flavivírus detetados em *Culex theileri*; CxunFV, Flavivírus detetados em *Culex univittatus*; OccaFV, Flavivírus detetados em *Ochlerotatus caspius*; WNV, vírus *West Nile*; DENV1, vírus Dengue serotipo 1; DENV2, vírus Dengue serotipo 2; DENV4, vírus Dengue serotipo 4.

... PREVENDO A POSSIBILIDADE DA EMERGÊNCIA DE AGENTES INFECIOSOS PARA OS QUAIS FORAM IDENTIFICADOS VETORES COMPETENTES...

Quadro 2 – Agentes etiológicos transmitidos por ixodídeos presentes ou em risco de emergir em Portugal

Agente patogénico	Doença	Espécie de ixodídeo
<i>Anaplasma phagocytophilum</i>	Anaplasmose humana	<i>Ixodes ricinus</i> , <i>I. ventralloii</i>
<i>Babesia divergens</i>	Babesiose	<i>Ixodes spp.</i>
<i>Borrelia burgdorferi s.s.</i>	Borreliose de Lyme	<i>Ixodes ricinus</i>
<i>B. garinii</i>		
<i>B. afzelii</i>		
<i>B. valaisiana</i>		
<i>B. lusitaniae</i>		
<i>B. turdi</i>	—	
<i>Coxiella burnetii</i>	Febre Q	Várias espécies
<i>Francisella tularensis</i>	Tularemia	Várias entre as quais <i>Ixodes ricinus</i> , <i>Dermacentor reticulatus</i>
<i>Rickettsia aeschlimannii</i>	Sem denominação	<i>Hyalomma marginatum</i>
<i>R. conorii</i>	Febre escaro nodular	<i>Rhipicephalus sanguineus</i>
<i>R. helvetica</i>	Sem denominação	<i>Ixodes ricinus</i>
<i>R. massillae</i>	Sem denominação	<i>Rhipicephalus sanguineus</i>
<i>R. monacensis</i>	Sem denominação	<i>Ixodes ricinus</i>
<i>R. sibirica mongolitimonae</i>	LAR*	<i>Hyalomma sp.</i> , <i>Rhipicephalus pusillus</i>
<i>R. slovaca</i>	TIBOLA [†]	<i>Dermacentor marginatus</i> , <i>D. reticulatus</i>
Vírus da Febre Hemorrágica Crimeia-Congo	Febre hemorrágica	<i>Hyalomma marginatum</i> , <i>Haemaphysalis punctata</i> , <i>Ixodes ricinus</i> , <i>Dermacentor spp.</i> , <i>Rhipicephalus spp.</i>
Vírus Eyach	Sem denominação	<i>Ixodes ricinus</i> , <i>Ixodes ventralloii</i>
Vírus TBE	Encefalite	<i>Ixodes ricinus</i> , <i>Haemaphysalis punctata</i>

* LAR - Lymphangitis-associated rickettsiosis; [†]TIBOLA - Tick-borne lymphadenopathy

... COMO ACONTECEU NESTES 2 CASOS!!!

CASO CLÍNICO / CLINICAL CASE

Infecção por vírus West Nile (Flavivírus) em Portugal

Considerações acerca de um caso clínico de síndrome febril com exantema

West Nile virus (Flavivirus) infection in Portugal

Considerations about a clinical case with febrile syndrome and rash

M. J. Alves¹ / J. M. D. Paços² / T. Luzz¹ / F. Amaro¹ / L. Zê-Zê³ / H. Oeirão¹

Centro de Estudos de Vetores e Doenças Infecciosas Dr. Francisco Cambouraz / Instituto Nacional de Saúde Dr. Ricardo Jorge
¹ Centro Hospitalar de Setúbal, Hospital S. Bernardo EPE

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/ Resumo

O vírus West Nile (WN) é um flavivírus transmitido por mosquitos e agente etiológico de febre e de doença neuroinvasiva. O vírus WN mantém-se na natureza em ciclos enzoóticos que envolvem mosquitos ornitofílicos, como vetores primários, e algumas espécies de aves como reservatório primário. A sua presença em Portugal é conhecida, surgindo esporadicamente alguns casos de infeção em equinos e humanos. Em 2010 foi identificado um caso humano na região sul de Portugal, tendo sido o único caso humano detectado em toda a época de actividade de mosquitos nesse ano.

Neste caso a paciente apresentava quadro febril com hiperpirexia muito irregular, por vezes com calafrios e picos de febre superiores a 39°C, cefaleias, mialgias, adinamia e astenia acentuada, adenomegalias volumosas e dolorosas na região cervical, assim como exantema eritematoso difuso com maior expressão no tronco. Os exames laboratoriais identificaram seroconversão de anticorpos IgM contra o vírus West Nile.

Palavras-chave: vírus West Nile; síndrome febril; zoonoses.

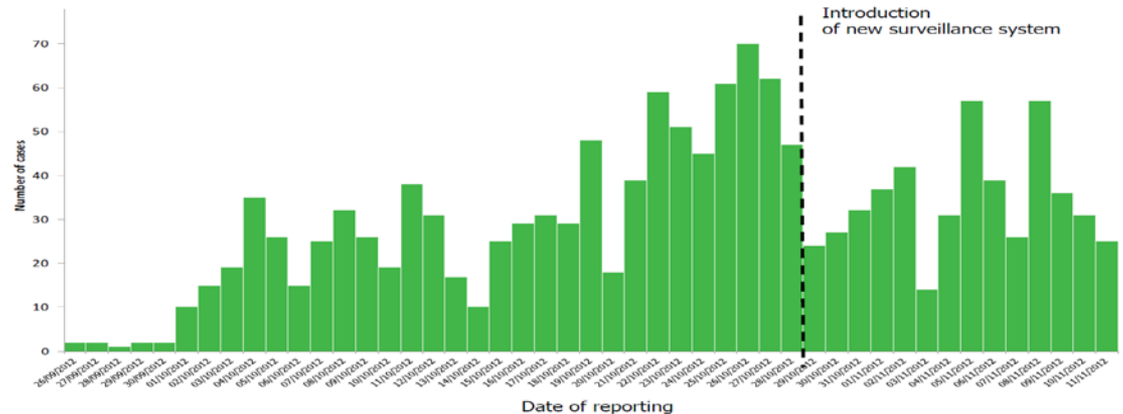


RAPID RISK ASSESSMENT

Update on autochthonous dengue cases in Madeira, Portugal

20 November 2012

Figure 1. Distribution of probable and confirmed dengue cases, by date of reporting, 26 September–11 November 2012, Madeira (n=1357)



A RESPONSABILIDADE DE ESTARMOS INSERIDOS NO SISTEMA NACIONAL DE VIGILÂNCIA DE VÍRUS RESPIRATÓRIOS

Instituto Nacional de Saúde
Doutor Ricardo Jorge

PORTUGAL

Boletim de Vigilância Epidemiológica da Gripe

Época 2015/2016

Semana 18 | 02 - 08 maio 2016

NOVA SÉRIE

Resumo

**Atividade gripal esporádica
Tendência estável**

Sumário

- Vigilância clínica**
Taxa de incidência de SG
- Vigilância laboratorial**
Diagnóstico do vírus da gripe e outros vírus respiratórios
Caraterização do vírus da gripe
- Severidade**
Internamentos por gripe em UCI
- Impacte**
Mortalidade por todas as causas
- Monitorização da temperatura ambiente, taxa de incidência de SG e mortalidade**
- Situação internacional**

- A taxa de incidência de síndrome gripal (SG) foi de 3,4 por 100.000 habitantes.
- Na semana 18/2016 foram identificados 14 casos de gripe pela Rede Portuguesa de Laboratórios para o Diagnóstico da Gripe. Os vírus da gripe detetados são predominantemente do tipo B.
- Os vírus da gripe circulantes são na sua maioria semelhantes aos vírus contemplados na vacina antigripal da época 2015/2016.
- Não foi admitido nenhum novo caso de gripe nas 20 UCI que reportaram informação.
- Mortalidade observada por todas as causas com valores de acordo com o esperado.
- O valor médio da temperatura mínima do ar, na semana de 2 a 8 de maio de 2016 foi de 8.1°C (valor próximo do normal).
- Na semana 17/2016 a maioria dos países referiu atividade gripal esporádica com tendência decrescente. Predomínio do vírus do tipo B.

Nota metodológica

ISSN: 2183-7392

Data de publicação: 12/05/2016

Dados disponíveis à data da publicação passíveis de alterações em edições posteriores.

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Vigilância laboratorial

Diagnóstico do vírus da gripe e outros vírus respiratórios

HOSPITAIS | REDE PORTUGUESA DE LABORATÓRIOS PARA O DIAGNÓSTICO DA GRIPES

Diagnóstico do vírus da gripe

A Rede Portuguesa de Laboratórios para o Diagnóstico da Gripe conta na época de 2015/2016 com a participação de 16 laboratórios, localizados em hospitais do continente e regiões autónomas da Madeira e dos Açores, assegurando a deteção e caracterização dos vírus da gripe e outros vírus respiratórios que podem estar associados a casos de infeção respiratória grave.

Desde o início da época 2015/2016, os laboratórios da Rede notificaram 6.748 casos de SG, dos quais 1.244 foram positivos para o vírus da gripe [914 vírus A(H1)pdm09, 177 vírus do tipo B, 21 vírus A(H3) e 125 vírus do tipo A não subtipados]. Nas últimas semanas foram detetados maioritariamente vírus da gripe do tipo B. Foram também identificados outros agentes respiratórios em 1.897 casos de SG sendo o vírus sincicial respiratório (RSV) o detetado com maior frequência. Nas últimas semanas verificou-se uma tendência decrescente no número de vírus respiratórios identificados.

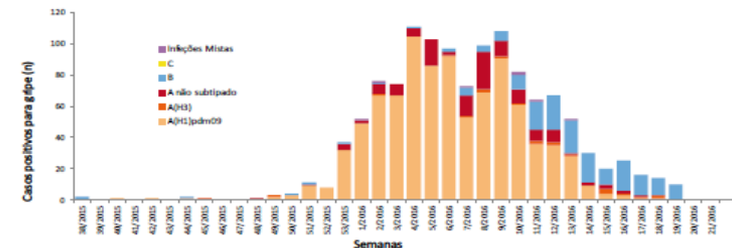


Figura 8 — Distribuição semanal de casos positivos para o vírus da gripe detetados na época 2015/2016.

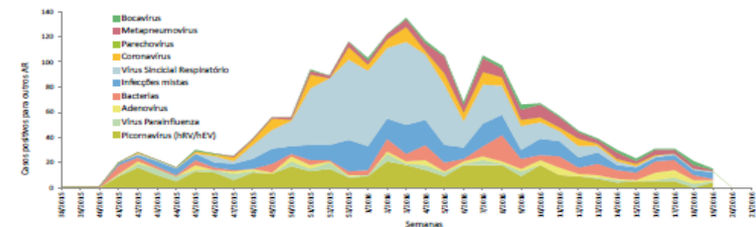


Figura 9 — Distribuição semanal de casos positivos para outros agentes respiratórios (AR) detetados na época 2015/2016.

A EXTREMA IMPORTÂNCIA DE ESTARMOS INTEGRADOS EM PROJETOS EUROPEUS

NEWS

Early influenza vaccine effectiveness results 2015-16: I-MOVE multicentre case-control study

E Kissling¹, M Valenciano¹

1. EpiConcept, Paris, France

Correspondence: Esther Kissling (e.kissling@epiconcept.fr)

Citation style for this article:

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Article published on 11 February 2016

Instituto Nacional de Saúde Doutor Ricardo Jorge, IP

ESTUDO DA EFETIVIDADE DA VACINA ANTIGRI PAL

EM CONTEXTO HOSPITALAR

2015-2016



CENTRO HOSPITALAR DE LISBOA CENTRAL, IPE

Centro Hospitalar de Setúbal Hospital de São Bernardo Hospital Ortopédico Sant'ago do Outão

I-MOVE

O Projeto EVA Hospital 2015-2016

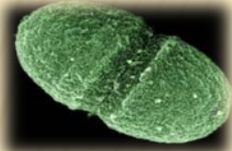
No âmbito do projeto I-MOVE+, está a ser implementado o estudo “Hospital-based test negative case control studies to measure seasonal influenza vaccine effectiveness against influenza laboratory confirmed SARI hospitalisation among the elderly across the European Union and European Economic Area Member States”, cujo objetivo é estimar a efetividade da vacina antigripal na prevenção de casos de gripe severos na população com 65 e mais anos.

Participam neste estudo 11 países (Figura 3), constituindo uma rede de 23 hospitais.

Portugal, participa pela primeira vez nesta rede hospitalar, contando com a colaboração do Centro Hospitalar Lisboa Central (CHLC) e Hospital São Bernardo (Centro Hospitalar de Setúbal), na implementação de um protocolo comum com um desenho do tipo caso-controlo teste negativo, selecionando casos graves de gripe (Severe Acute Respiratory Infection-SARI).

Nesta época 2015-2016, a fase de seleção de doentes com SARI iniciou-se na semana 51/2015 em CHLC e na semana 1/2016 em CHS.

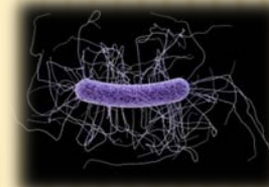
MICROORGANISMOS “ALERTA” E “PROBLEMA”: A PARTICIPAÇÃO NA REDE DE VIGILÂNCIA DA DGS E DO ECDC



ENTEROCOCCUS
- RESISTENTE À VANCOMICINA (VRE)



NEISSERIA GONORRHOEAE



CLOSTRIDIUM DIFFICILE



ENTEROBACTERIACEAE
- RESISTENTES AOS CARBAPENEMES



PSEUDOMONAS AERUGINOSA



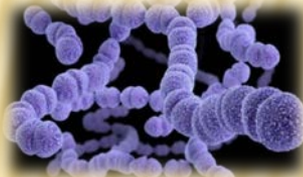
SHIGELLA



STAPHYLOCOCCUS AUREUS
METHICILLIN-RESISTANT (MRSA)



SALMONELLA TYPHI



STREPTOCOCCUS PNEUMONIAE



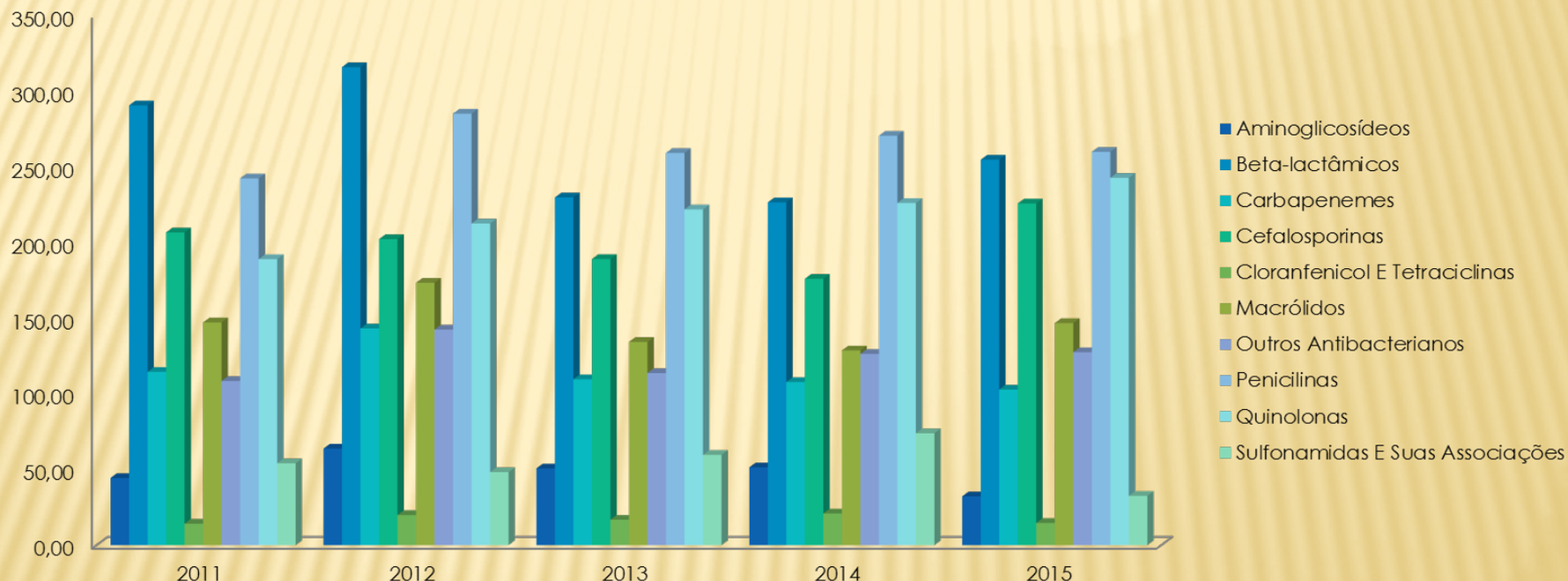
CAMPYLOBACTER



ENTEROBACTERIACEAE
- PRODUTORAS DE β -LACTAMASES

ANÁLISE DO CONSUMO DE ANTIBIÓTICOS

Famílias Antibióticas 2011-2015



Centro Hospitalar de Setúbal
Hospital de São Bernardo
Hospital Ortopédico Sant'Iago do Outão

- Acréscimo de 2% no total de Antibióticos consumidos de 2014 para 2015;
- 4 grandes famílias: **Beta-lactâmicos, Cefalosporinas, Penicilinas e Quinolonas;**

ELABORANDO E DIVULGANDO PERIODICAMENTE A CARTA MICROBIOLÓGICA DO CHS

Centro Hospitalar de Setubal, EPE
Grupo Coordenador Local de Controlo de
Infeção e de Prevenção de Resistências aos
Antimicrobianos (GCLCIPRA)

**Perfil de sensibilidade (%) das
principais bactérias isoladas em 2015**

		Bactérias Gram -						Bactérias Gram +				
		<i>Escherichia coli</i>	<i>Klebsiella pneumoniae</i>	<i>Proteus mirabilis</i>	<i>Pseudomonas aeruginosa</i>	<i>Acinetobacter baumannii</i>	<i>Acinetobacter baumannii complex</i>	<i>Haemophilus influenzae</i>	<i>Staphylococcus aureus</i>	<i>Enterococcus faecalis</i>	<i>Enterococcus faecium</i>	<i>Streptococcus pneumoniae</i>
Nº amostras		1315	266	178	252	4	25	9	555	197	21	21
β-lactâmicos	Oxacilina								46			
	Ampicilina	42		62				88		100	24	100
	Amoxicilina/Ác.Clavulânico	81	69	93				100				
	Cefotaxima	89	72	95				100				100
	Ceftazidima	89	70	95	79	75	38					
	Piperacilina/Tazobactam	93	72	99	78	75	41					
	Imipenem; Meropenem	100	98	100	78	75	50	100		100	24	100
Aminoglicosidos	Gentamicina	87	80	88	86	75	54	100	95			
	Gentamicina_HC								72	65		
	Amikacina	95	89	97	88	100	80					
Quinolonas	Ciprofloxacina	72	70	66	69	75	46	100	39	82	6	100
Macrólidos	Eritromicina											74
Glicopéptidos	Vancomicina							100	100	95	100	
Sulfamidas	Cotrimoxazol	65	73	65		75	52	100	98			85

■ < 50% de estirpes sensíveis
■ Entre 50 a 85% de estirpes sensíveis
■ >85% de estirpes sensíveis

CARACTERIZAÇÃO DA CASUÍSTICA DE AMBULATÓRIO DO SDI: TOTAL DE DOENTES: 5676 (+ 446) EM JANEIRO 2016

× HIV

- + Total- 2753 (+ 156)
 - × Ativos: 1216 (+ 156)
 - × ADMITIDOS DE NOVO: 65 (-4)
 - * HIV1: 1143 (+ 207)
 - * HIV2: 23 (+ 3)
 - * Presos: 50 (=)
 - × Falecidos Total: 735 (+17)
 - * Em 2015: 17
 - × Refratários: 667 (+82)
 - × Transferidos: 135 (-7)

× Hepatites víricas (Mono-infetados)

- + Total: 992 (+64)
 - × Ativos: 521
 - × ADMITIDOS DE NOVO: 98
 - × Presos: 26
- + Falecidos: 156
 - × Em 2015: 1
- + Não ativos: 310
- + Transferidos: 5

DUAS REALIDADES MUITO DISTINTAS... MAS COM ALGUNS PONTOS EM COMUM...

× HIV

+ Cenário Presente

- × Refratários: 35,4%
- × Doentes em TARc: 920
- × % de D. em TT
 - * 75,6% (D. Ativos)
 - * 48,9% (D. Existentes)
- × % de D. c/ CV < 20
 - * 80% (D. sob TARc)
 - * 60% (D. Ativos)
 - * 39,1% (D. Existentes)

+ Cenário “Real”

- × Total de Infetados: 3500 (+ 750)
 - * % de D. em TT: 26,3%
 - * % de D. c/ CV < 20: 21%

× Hepatites (Mono-infecção)

- + Refratários: 37,3%

× Exercício Teórico

- + 95% c/ HCV= 800 Doentes
 - × 25%: Ac. HCV + c/ PCV -
 - × 25% já anteriormente Tratados
 - × 120 c/ DAA
- + Admite-se existirem presentemente: 280 Doentes ainda para tratar

× Se a prevalência e o n° p/ diagnosticar for idêntica ao HIV: + 300 Doentes

- + Se 25% tiverem Ac + e CV -
 - × Ficam: 225 Doentes

× Total p/ Tratar: 405 doentes(?)

OS RESULTADOS DE DUAS IMPORTANTES INICIATIVAS

× Consulta do Viajante

+ Dos 48 que aceitaram
fazer o teste rápido p/
HIV

× 1 Positivo (2%)

× Projeto da Unidade Móvel do GAT

+ Doentes referenciados
em 2015-2016: 25

× Mantêm-se em seguimento:
11

★ HIV positivos: 3

★ HCV positivos: 9

× Altas e Transferências: 3

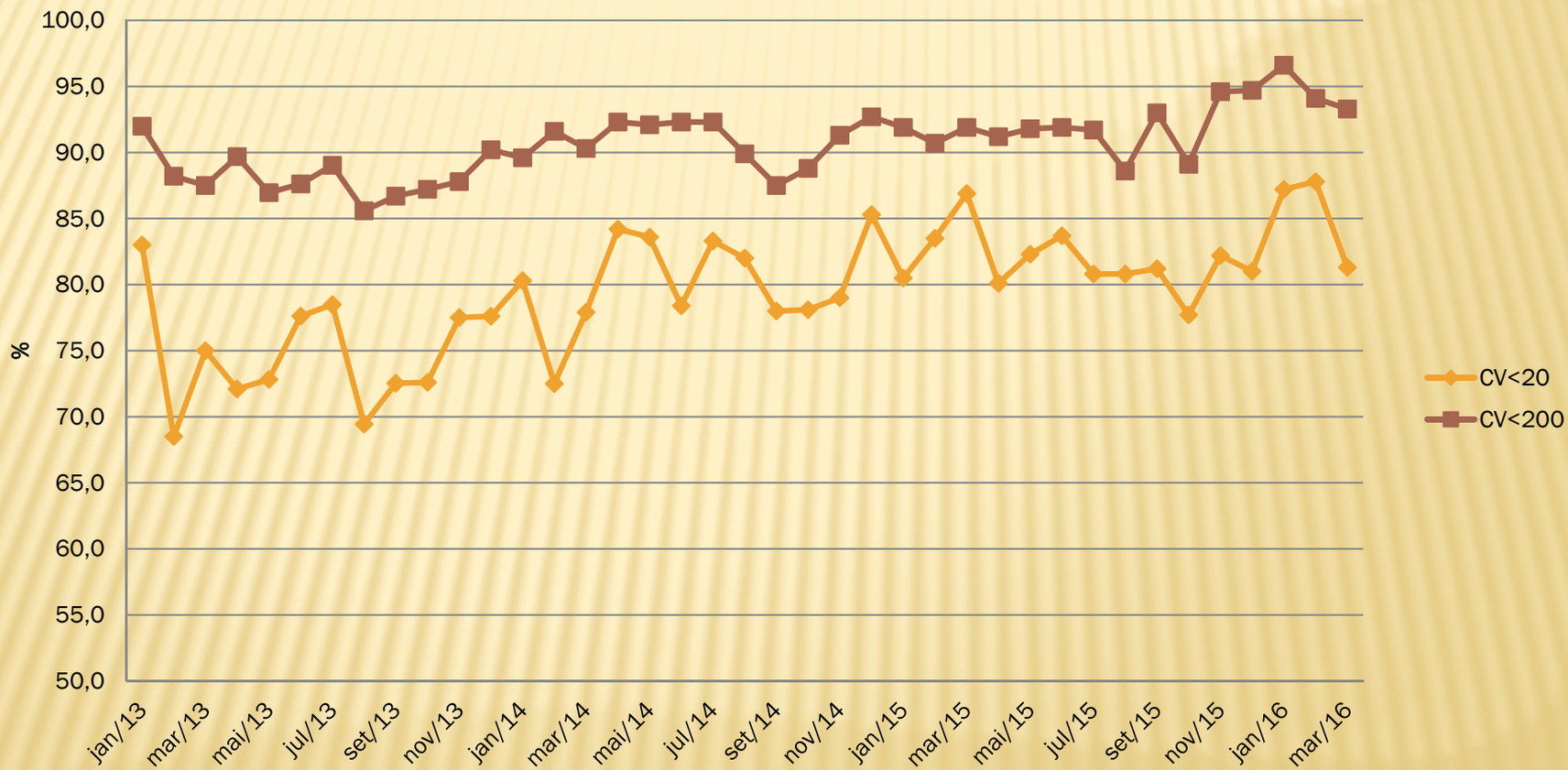
× Abandonaram: 11 (44%)

★ HIV: 1

★ HCV: 10

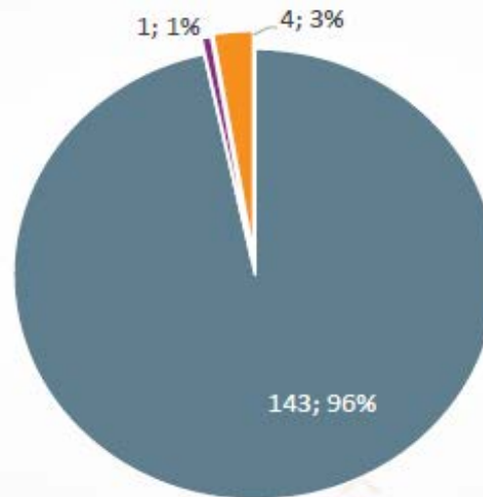
A NECESSIDADE DE MONITORIZARMOS EM PERMANÊNCIA E ADEQUADAMENTE A NOSSA ATUAÇÃO

Doentes em ART com CV suprimidas



RESULTADOS DA NOSSA EXPERIÊNCIA NO TRATAMENTO DA HEPATITE C C/ OS NOVOS FÁRMACOS

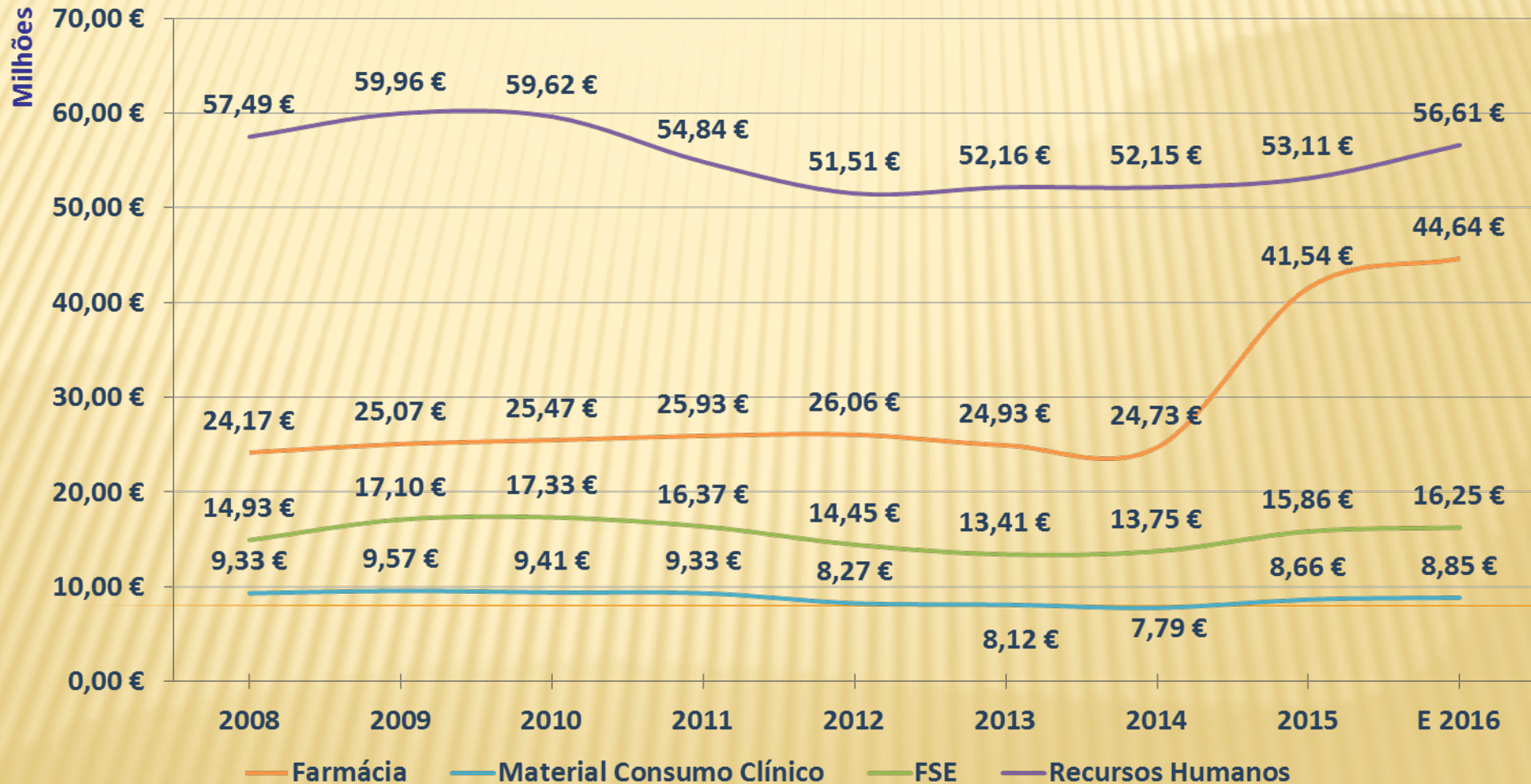
RESPOSTA VIROLÓGICA SUSTENDADA



■ Doentes com RVS ■ Doentes sem RVS ■ Aguardam avaliação analítica

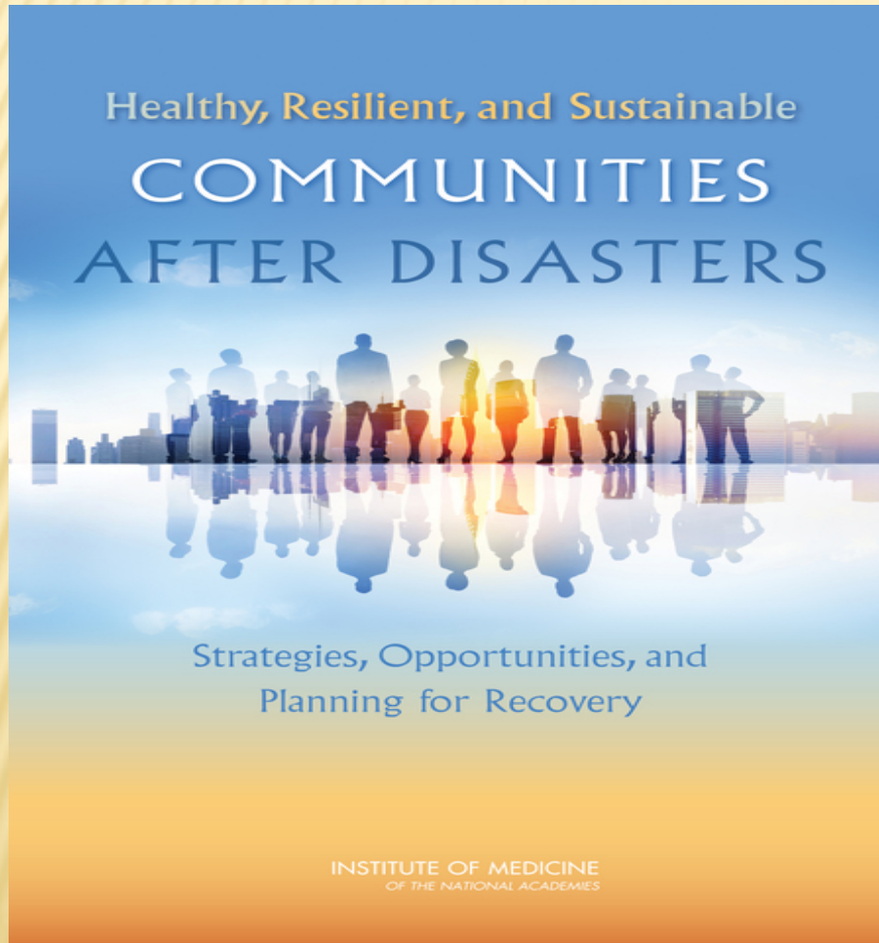


Evolução de Custos



VI)- CONCLUSÕES

TER EM CONSIDERAÇÃO AS NOVAS REALIDADES EPIDEMIOLÓGICAS ...



World Health Organization
REGIONAL OFFICE FOR **Europe**

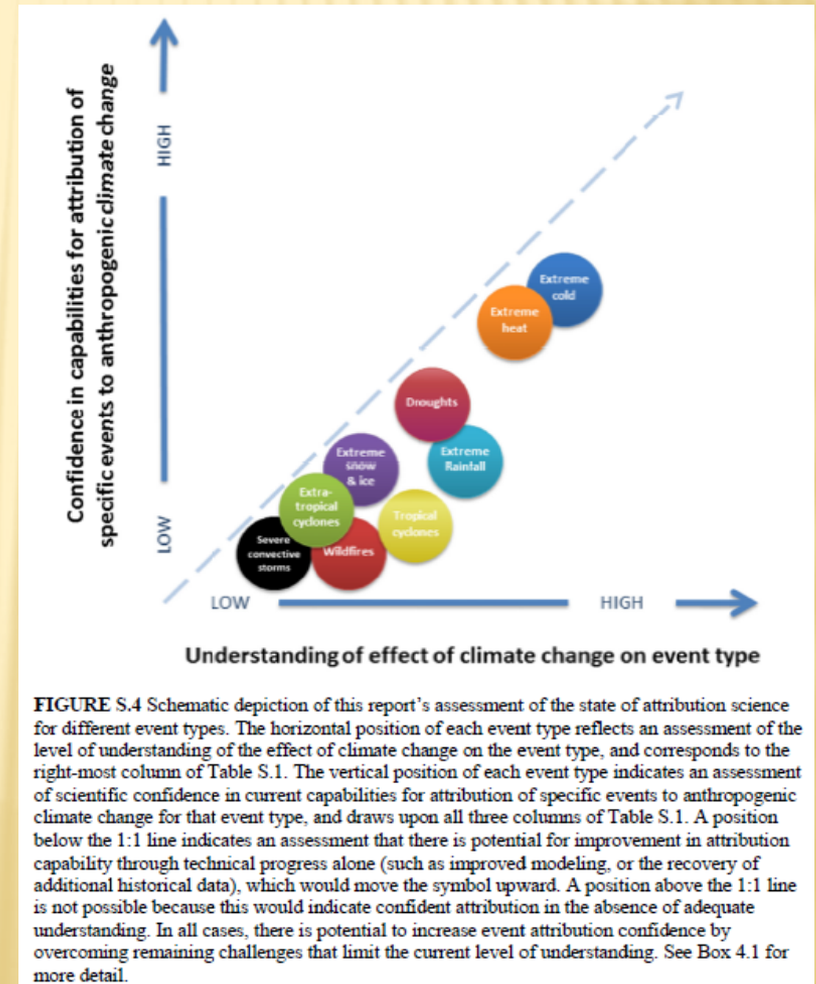
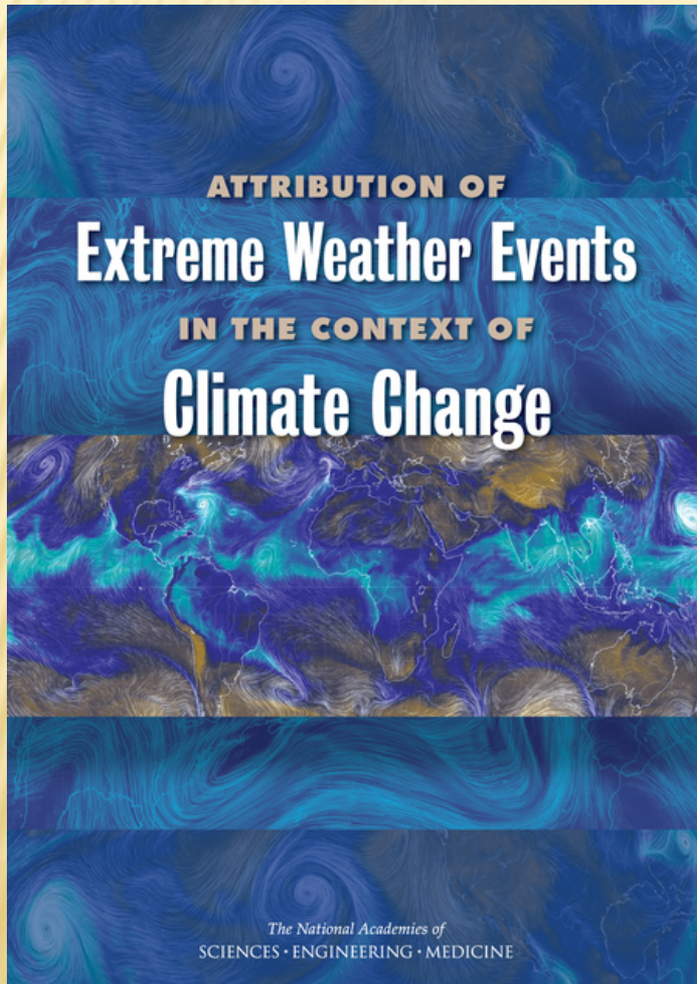
Health planning for large public events

Rio Olympics could spark 'full blown global health disaster', say Harvard scientists

Zika warning is latest in string of troubles for 2016 Games hosts

The complex block contains the WHO logo and the text 'World Health Organization REGIONAL OFFICE FOR Europe'. Below this is the title 'Health planning for large public events'. It features two images: one showing people in yellow hazmat suits, and another showing a hand holding a torch over a chaotic scene with a bicycle and other debris. At the bottom, there is a headline: 'Rio Olympics could spark 'full blown global health disaster', say Harvard scientists' and a sub-headline: 'Zika warning is latest in string of troubles for 2016 Games hosts'.

... TAIS COMO OS EVENTOS METEOROLÓGICOS EXTREMOS...



... E TAMBÉM O BIOTERRORISMO, OS GRANDES MOVIMENTOS MIGRATÓRIOS E AS VIAGENS AÉREAS!!!



A DECADE OF BIODEFENCE

IMMIGRATION ROUTES INTO THE EU BY LAND AND SEA

Most migrants enter the EU through international airports, most of those living in the EU illegally originally entered with valid documents but then overstayed a visa. Many others though enter via land and sea routes. The estimated numbers of those who entered in 2012 are shown below.



SOURCE: Frontex Annual Risk Analysis 2013 | Designed by Inez Torre/CNN



FIGURE A6-2 The global aviation network. Lines show direct links between airports, and the colour indicates passenger capacity in people per day (thousands [red]; hundreds [yellow]; tens [blue]). Routes linking regions at similar latitudes (in the northern or southern hemisphere) represent pathways that pathogens can move along to reach novel regions. Notably, air traffic to most places in Africa, regions of South America, and parts of central Asia is low. If travel increases in these regions, additional introductions of vector-borne pathogens are probable.
SOURCE: Adapted from Hufnagal et al., 2004.

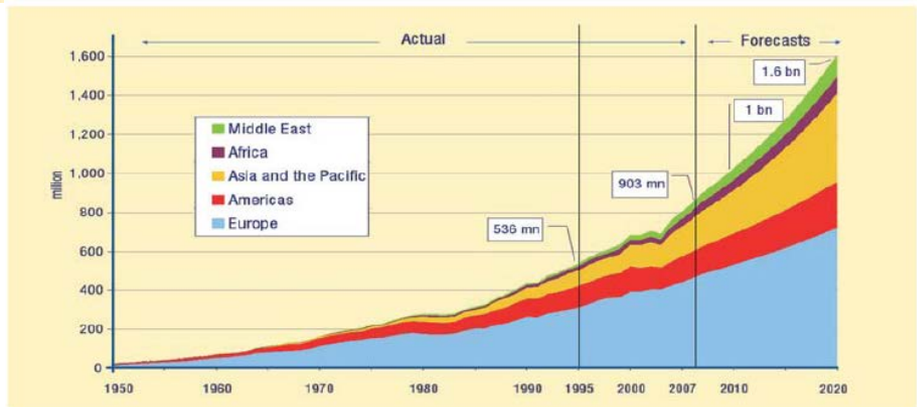


FIGURE WO-4 International tourist arrivals by region (in millions), 1950-2020.
SOURCE: Reprinted with permission from the UNWTO (2008).

TRATA-SE EFETIVAMENTE DE UMA QUESTÃO DE SEGURANÇA!!!

The screenshot shows the White House website with a navigation bar at the top containing links for President, Vice President, First Lady, Mrs. Cheney, and News. Below the navigation bar is a search box and a "Search" button. The main content area features a banner for "WAR ON TERROR PROJECT BIOSHIELD" with the subtitle "PROGRESS IN THE WAR ON TERROR". Below the banner is a photo of President George W. Bush signing the Project BioShield Act of 2004, surrounded by other officials. To the left of the photo is a sidebar with "IN FOCUS" categories and "News" items. To the right is another sidebar with "SPEECHES & NEWS RELEASES" and "ASK THE WHITE HOUSE" sections.



Public Policy



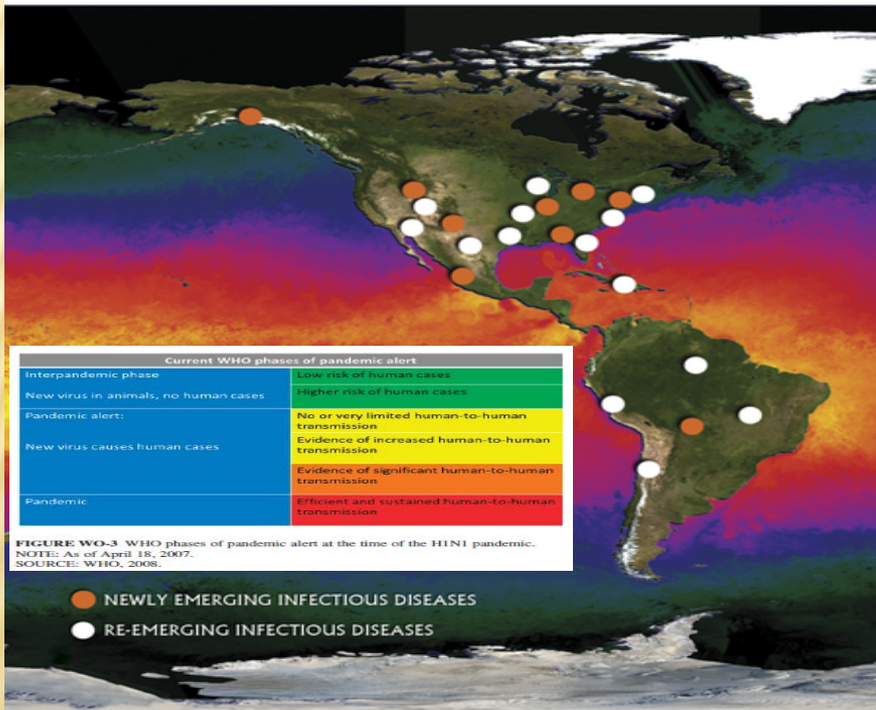
Global health security: the wider lessons from the west African Ebola virus disease epidemic

David L. Heymann, Lincoln Chen, Kefau Takemi, David P. Fidler, Jordan W. Tappero, Michael Thomas, Thomas A. Keryon, Thomas R. Frieden, Derek Yach, Sania Nishtar, Alex Katsche, Piero L. Ollano, Peter Harby, Elis Torredé, Lawrence O. Gostin, Margaret Ndomondo-Sigonda, Daniel Carpenter, Simon Bushoro, Louis Lillywhite, Bhimsen Deukota, Khalid Koser, Rob Yates, Rana S. Dhillon, Ravi P. Raman-Ebya

The Ebola virus disease outbreak in West Africa was unprecedented in both its scale and impact. Out of this human calamity has come renewed attention to global health security—its definition, meaning, and the practical implications for programmes and policy. For example, how does a government begin to strengthen its core public health capacities, as demanded by the International Health Regulations? What counts as a global health security concern? In the context of the governance of global health, including WHO reform, it will be important to distil lessons learned from the Ebola outbreak. *The Lancet* invited a group of respected global health practitioners to reflect on these lessons, to explore the idea of global health security, and to offer suggestions for next steps. Their contributions describe some of the major threats to individual and collective human health, as well as the values and recommendations that should be considered to counteract such threats in the future. Many different perspectives are proposed. Their common goal is a more sustainable and resilient society for human health and wellbeing.

A NECESSIDADE DE UMA PERMANENTE E RÁPIDA VIGILÂNCIA ATRAVÉS DE MEIOS EFICAZES...

THE INFLUENCE OF GLOBAL ENVIRONMENTAL CHANGE ON INFECTIOUS DISEASE DYNAMICS



Current WHO phases of pandemic alert	
Interpandemic phase	Low risk of human cases
New virus in animals, no human cases	Higher risk of human cases
Pandemic alert:	No or very limited human-to-human transmission
New virus causes human cases	Evidence of increased human-to-human transmission
	Evidence of significant human-to-human transmission
Pandemic	Efficient and sustained human-to-human transmission

FIGURE WO-3 WHO phases of pandemic alert at the time of the H1N1 pandemic.
NOTE: As of April 18, 2007.
SOURCE: WHO, 2008.

- NEWLY EMERGING INFECTIOUS DISEASES
- RE-EMERGING INFECTIOUS DISEASES

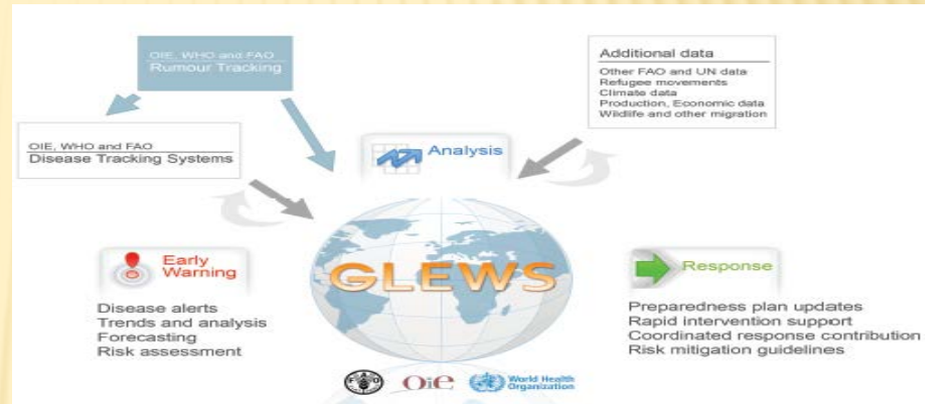


FIGURE 5-1 Global Early Warning and Response System (GLEWS) for Major Animal Diseases, including Zoonoses.
SOURCE: OIE (2009).

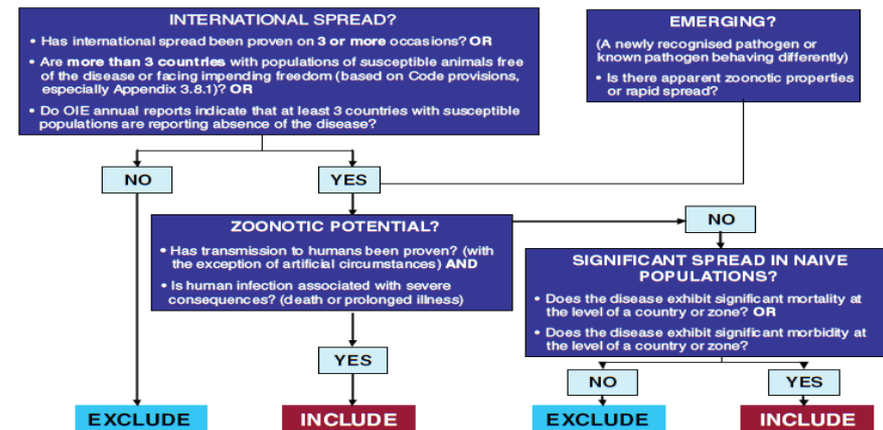


FIGURE 5-3 OIE's disease notification criteria.
SOURCE: OIE (2008).

... E DE SABER UNIR ESFORÇOS!!!

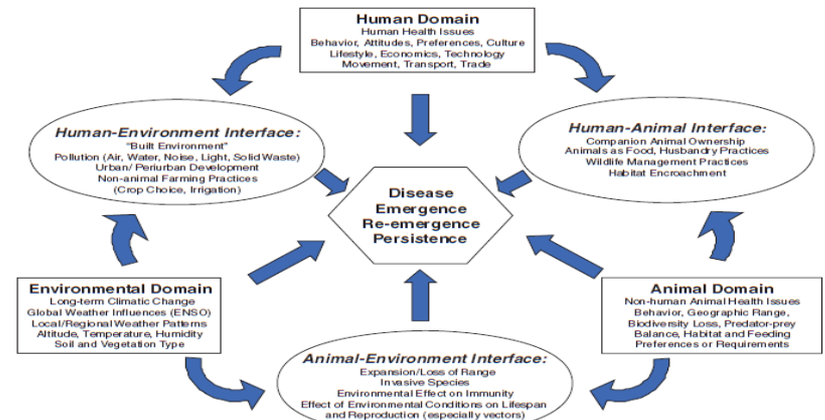
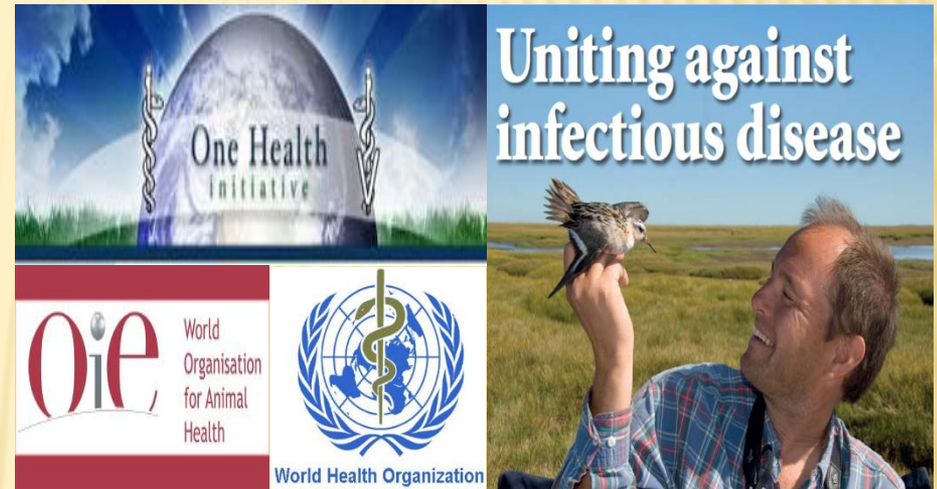
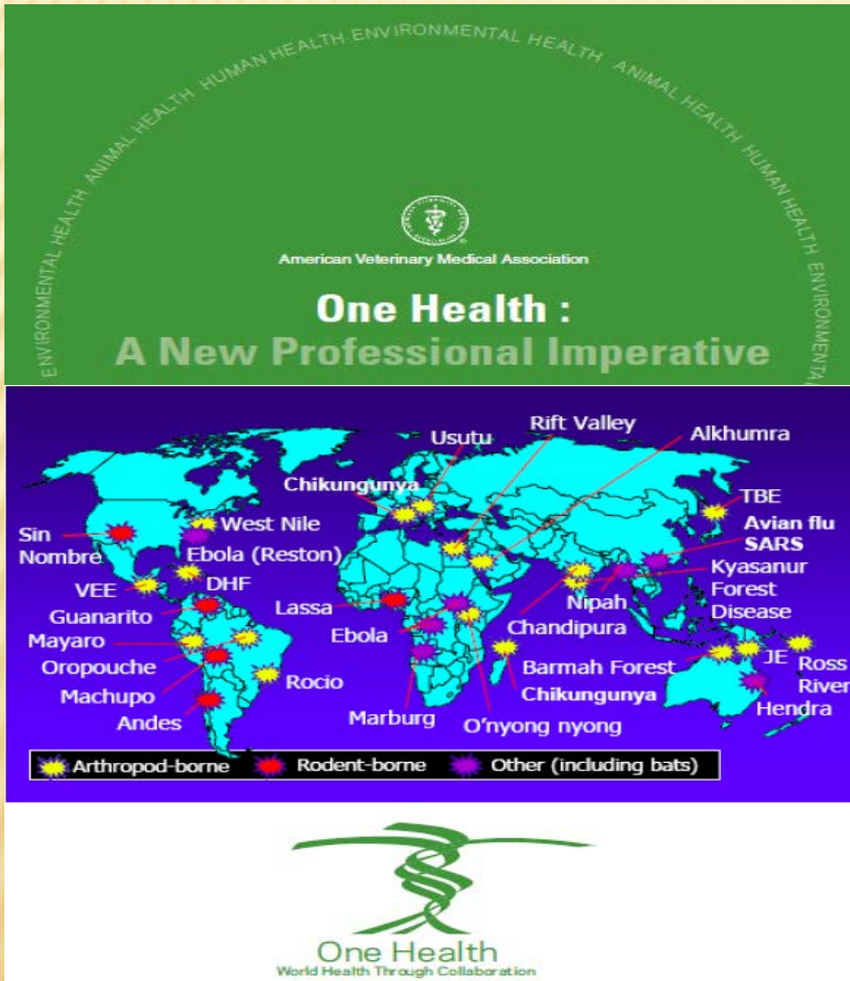
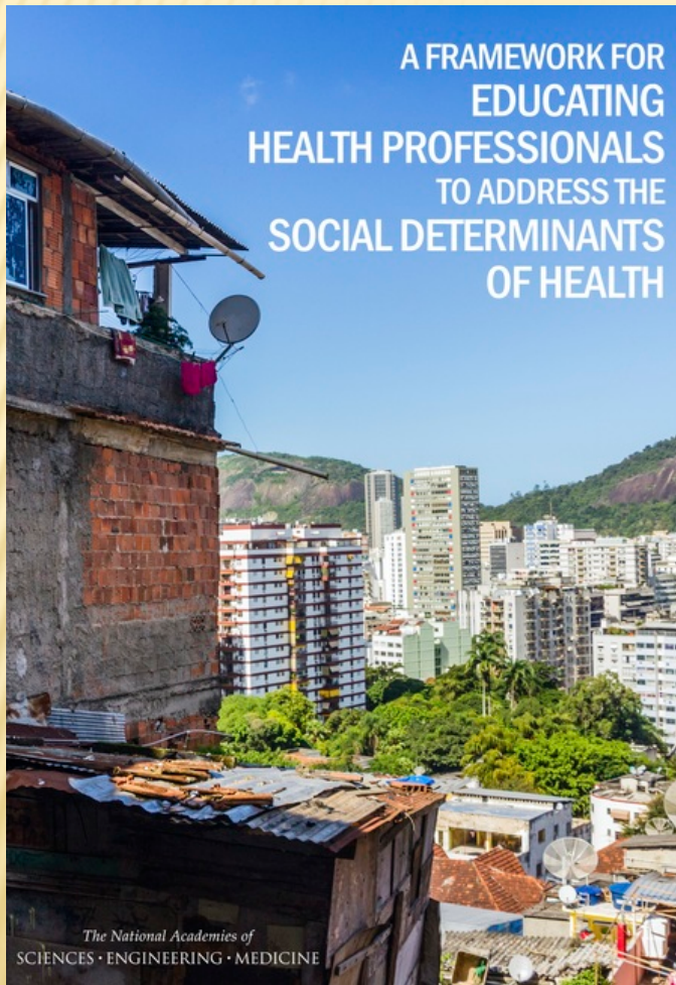


FIGURE 3-1 Overview of the driver-pathogen interactions that contribute to the emergence of infectious zoonotic diseases. SOURCE: Treadwell (2008).

É NECESSÁRIO FAZER POIS UM GRANDE ESFORÇO PARA A EDUCAÇÃO CÍVICA DOS PROFISSIONAIS E DOS CIDADÃOS...



AS AJUDAS QUE NECESSITAMOS...

SUPPLEMENT ARTICLE

Use of Electronic Health Records and Clinical Decision Support Systems for Antimicrobial Stewardship

Graeme N. Forrest,¹ Trevor C. Van Schooneveld,² Ravina Kullar,³ Lucas T. Schulz,⁴ Phu Duong,⁵ and Michael Postelnick⁶

¹Division of Infectious Diseases, Portland Veterans Affairs Medical Center, Portland, Oregon; ²University of Nebraska Medical Center, Omaha; ³Global Medical Affairs, Cubist Pharmaceuticals, Lexington, Massachusetts; ⁴University of Wisconsin Hospital and Clinics, Madison; and ⁵Northwestern Memorial Hospital, Chicago, Illinois

Electronic health records (EHRs) and clinical decision support systems (CDSSs) have the potential to enhance antimicrobial stewardship. Numerous EHRs and CDSSs are available and have the potential to enable all clinicians and antimicrobial stewardship programs (ASPs) to more efficiently review pharmacy, microbiology, and clinical data. Literature evaluating the impact of EHRs and CDSSs on patient outcomes is lacking, although EHRs with integrated CDSSs have demonstrated improvements in clinical and economic outcomes. Both technologies can be used to enhance existing ASPs and their implementation of core ASP strategies. Resolution of administrative, legal, and technical issues will enhance the acceptance and impact of these systems. EHR systems will increase in value when manufacturers include integrated ASP tools and CDSSs that do not require extensive commitment of information technology resources. Further research is needed to determine the true impact of current systems on ASP and the ultimate goal of improved patient outcomes through optimized antimicrobial use.

Keywords. antimicrobial stewardship; clinical decision support system; electronic health record.

Table 2. Clinical Decision Support Systems and Patient Outcomes

Reference Number	Study Design	Software	Setting	Results	Notes
[9]	Pre-Post	TheraDoc	ICU	Significant declines in antibiotic susceptibility mismatches, duration of excess drug doses, and orders for antibiotics to which the patient was allergic ($P < .01$). Also had a 70% reduction in ADE ($P = .018$).	No differences in mortality between groups
[10]	Prospective	TheraDoc	Inpatient	22.8% decline in antibiotic use, a \$70 per-patient decrease in antibiotic costs, a decline in antibiotic adverse events, and a decline in hospital mortality over a 7-year period (3.65% to 2.65%, $P < .001$).	Time period evaluated was from 1988 to 1994
[56]	Cluster randomized	TheraDoc	Community clinics	Antibiotic prescribing rate declined from 84.1 to 75.3 prescriptions per 100 person-years ($P = .03$). Also reduced inappropriate antibiotic prescribing, from 32% to 5% ($P = .03$).	Macrolides reduced 28%, cephalosporins 7%, and penicillins 6%
[57]	Pre-post	Unknown	PPRnet—outpatients	Inappropriate antibiotic use declined 0.6% for ARI and 16.6% for broad antibiotics in adults.	Modest effect
[58]	Prospective interventional	Unknown	PPRnet	Antibiotic use did not change (+1.57%), decrease in broad antibiotic use for ARI (−16%).	Decreased broad antibiotic use
[59]	Retrospective observational	Unknown	Veterans Affairs—outpatients	Increase in antibiotic usage (0.63 to 0.72, $P = .001$).	No effect seen targeting ARI antibiotics
[60]	Prospective	Local program	Outpatients	Overall antibiotic prescribing 39% vs non-CDSS of 43%. ARI was 54% vs 59%.	CDSS form only used in 6% of ARI visits
[61]	Prospective	TheraDoc	Pediatrics	59% reduction in erroneous antimicrobial use, 28% decline in excess dose-days. No change in ADE or susceptibility mismatches.	
[62]	Cluster-randomized study	TREAT	Inpatient	Better empiric antibiotic therapy (70% vs 57%, $P < .001$). Length of stay and costs (−12%) also reduced.	No impact on mortality
[63]	Survival analysis	TREAT	Inpatient, single center	The ITT group 180-day survival in the control group was 68% vs 71% in the intervention group ($P = .1$). In the PP analysis, the survival percentages were 68% vs 74% ($P = .04$).	Analysis of only 1 center of whole study that analyzed 30-day mortality
[64]	Prospective	Antibiograms	ICU	Increased susceptibility to imipenem (18.3%/year) and gentamicin (11.6%/year).	No clinical outcomes data.

Abbreviations: ADE, adverse drug event; ARI, acute respiratory infection; CDSS, clinical decision support system; ICU, intensive care unit; ITT, intent to treat; PP, per protocol; PPRnet, Practice Partners Research Network.

... COMO NESTE EXEMPLO...

BJCP British Journal of Clinical Pharmacology

Therapeutic drug monitoring of antimicrobials

Jason A. Roberts,^{1,4,5} Ross Norris,^{2,7,8} David L. Paterson^{3,6} & Jennifer H. Martin⁹

¹Burns, Trauma and Critical Care Research Centre, ²School of Pharmacy and ³Centre for Clinical Research, The University of Queensland, Brisbane, Queensland, Australia, ⁴Department of Intensive Care, ⁵Pharmacy Department and ⁶Department of Infectious Diseases, Royal Brisbane and Women's Hospital, Brisbane, Queensland, Australia, ⁷Australian Centre for Paediatric Pharmacokinetics, Mater Pharmacy Services, Brisbane, Queensland, Australia, ⁸School of Pharmacy, Griffith University, Gold Coast, Queensland, Australia and ⁹The University of Queensland School of Medicine Southside, Princess Alexandra Hospital, Woolloongabba, Brisbane, Queensland, Australia

Wong et al. *BMC Infectious Diseases* 2014, **14**:288
<http://www.biomedcentral.com/1471-2334/14/288>



REVIEW

Open Access

How do we use therapeutic drug monitoring to improve outcomes from severe infections in critically ill patients?

Gloria Wong^{1†}, Fekade Bruck Sime^{2,3†}, Jeffrey Lipman^{1,4} and Jason A Roberts^{1,2,4*}

Abstract

High mortality and morbidity rates associated with severe infections in the critically ill continue to be a significant issue for the healthcare system. In view of the diverse and unique pharmacokinetic profile of drugs in this patient population, there is increasing use of therapeutic drug monitoring (TDM) in attempt to optimize the exposure of antibiotics, improve clinical outcome and minimize the emergence of antibiotic resistance. Despite this, a beneficial clinical outcome for TDM of antibiotics has only been demonstrated for aminoglycosides in a general hospital patient population. Clinical outcome studies for other antibiotics remain elusive. Further, there is significant variability among institutions with respect to the practice of TDM including the selection of patients, sampling time for concentration monitoring, methodologies of antibiotic assay, selection of PK/PD targets as well as dose optimisation strategies. The aim of this paper is to review the available evidence relating to practices of antibiotic TDM, and describe how TDM can be applied to potentially improve outcomes from severe infections in the critically ill.

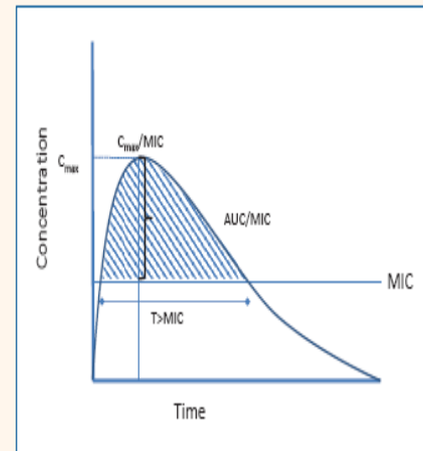
Keywords: TDM, Antibiotic, Pharmacokinetics, Pharmacodynamics

Comparative Effectiveness Review
Number 136

Effective Health Care Program

Pharmacokinetic/Pharmacodynamic Measures for Guiding Antibiotic Treatment for Hospital-Acquired Pneumonia Executive Summary

Figure A. Ratios related to the minimum inhibitory concentration of the organisms



AUC = antibiotic area under the curve; AUC/MIC = the ratio of the antibiotic area under the curve to the time above the minimum inhibitory concentration needed to inhibit microorganisms; C_{max} = the maximum serum concentration needed to inhibit microorganisms; C_{max}/MIC = ratio of maximum serum concentration (or peak) to the time above the minimum inhibitory concentration needed to inhibit microorganisms; MIC = minimum inhibitory concentration; T = time.

... A IMPLEMENTAÇÃO DE TECNOLOGIAS QUE PERMITAM ENCURTAR COM SEGURANÇA A DURAÇÃO DOS CURSOS DE TRATAMENTO ...

Schuetz et al. *BMC Medicine* 2011, 9:107
<http://www.biomedcentral.com/1741-7015/9/107>



REVIEW

Open Access

Procalcitonin for diagnosis of infection and guide to antibiotic decisions: past, present and future

Philipp Schuetz^{1*}, Werner Albrich² and Beat Mueller²

Abstract

There are a number of limitations to using conventional diagnostic markers for patients with clinical suspicion of infection. As a consequence, unnecessary and prolonged exposure to antimicrobial agents adversely affect patient outcomes, while inappropriate antibiotic therapy increases antibiotic resistance. A growing body of evidence supports the use of procalcitonin (PCT) to improve diagnosis of bacterial infections and to guide antibiotic therapy. For patients with upper and lower respiratory tract infection, post-operative infections and for severe sepsis patients in the intensive care unit, randomized-controlled trials have shown a benefit of using PCT algorithms to guide decisions about initiation and/or discontinuation of antibiotic therapy. For some other types of infections, observational studies have shown promising first results, but further intervention studies are needed before use of PCT in clinical routine can be recommended. The aim of this review is to summarize the current evidence for PCT in different infections and clinical settings, and discuss the reliability of this marker when used with validated diagnostic algorithms.

Cost-Effectiveness of Procalcitonin-Guided Antibiotic Therapy for Outpatient Management of Acute Respiratory Tract Infections in Adults

Constantinos I. Michaelidis, BA¹, Richard K. Zimmerman, MD, MPH², Mary Patricia Nowalk, PhD², Michael J. Fine, MD, MSc^{3,4}, and Kenneth J. Smith, MD, MS³

¹University of Pittsburgh School of Medicine, Pittsburgh, PA, USA; ²Department of Family Medicine and Clinical Epidemiology, University of Pittsburgh School of Medicine, Pittsburgh, PA, USA; ³Division of General Internal Medicine, University of Pittsburgh School of Medicine, Pittsburgh, PA, USA; ⁴Center for Health Equity Research and Promotion, VA Pittsburgh Healthcare System, Pittsburgh, PA, USA.

Observational studies

Intervention studies

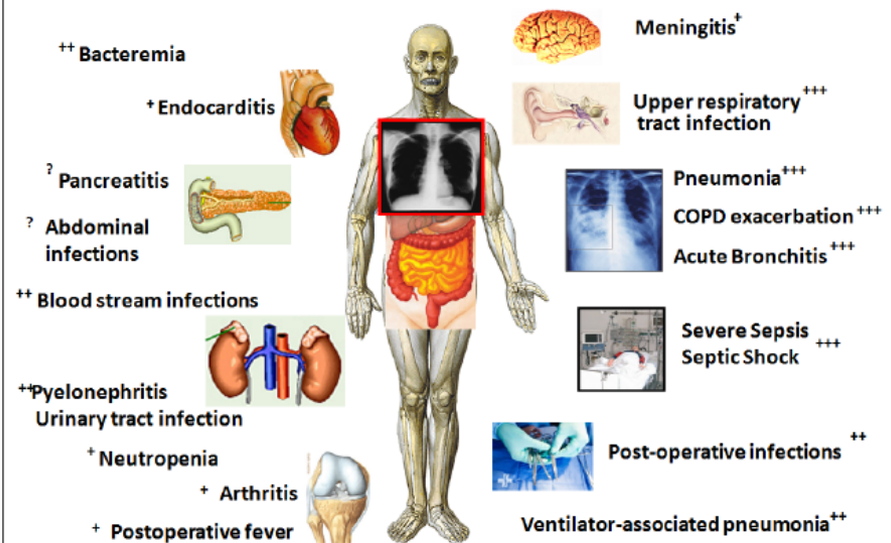


Figure 1 Available evidence concerning PCT in different infections derived from observational and randomized-controlled intervention studies. While for some infections, intervention studies have investigated benefit and harm of using PCT for antibiotic decisions (right side), for other infections only results from diagnostic (observation) studies are available with mixed results (left side). Abbreviations: PCT, procalcitonin. + moderate evidence in favor of PCT; ++ good evidence in favor of PCT; +++ strong evidence in favor of PCT; ? evidence in favor or against the use of PCT still undefined

AQUILO A ESTAMOS OBRIGADOS...



PATIENT SAFETY AND HEALTHCARE-ASSOCIATED INFECTIONS: *REPORT*
FROM THE
COMMISSION TO THE COUNCIL June 2014

2. ESTRUTURA DE GESTÃO E OPERACIONALIZAÇÃO DO PROGRAMA

Figura 1. Estrutura de gestão do Programa de Prevenção e Controlo de Infeções e de Resistência aos Antimicrobianos (PPCIRA)

Estrutura de gestão do PPCIRA



Fonte: PPCIRA / DGS / 2013

... A EXTREMA NECESSIDADE DE DISPOR CE MEIOS PARA GARANTIR OS FUNDAMENTAIS “LINKAGE” E O “RETENÇION TO CARE” ...

Linkage, Engagement, and Retention in HIV Care Among Vulnerable Populations Volume 21 Issue 4 September/October 2013

Perspective

Linkage, Engagement, and Retention in HIV Care Among Vulnerable Populations: “I’m Sick and Tired of Being Sick and Tired”

There are disparities in engagement and retention in HIV care and outcomes of care across segments of society. For example, HIV mortality rates remain markedly elevated among black women and men compared with their white counterparts. These differences reflect broader disparities across social, economic, and cultural lines. Improvement in engagement and retention in HIV care requires interventions that account for forces present in the socioecologic framework of health behaviors. Improvement in linkage to care at HIV testing is crucial to overall engagement and retention in care. Strategies for linkage to care at testing can help overcome many of the forces that result in failure to engage and remain in care by starting the patient on a solid path to clinical care. This article summarizes a presentation by Victoria A. Cargill, MD, MSCE, at the IAS–USA continuing education program held in New York, New York, in May 2013.

Keywords: HIV care disparities, linkage to care, socioecologic framework, engagement and retention in care, practitioner behavior

IAS–USA

Topics in Antiviral Medicine

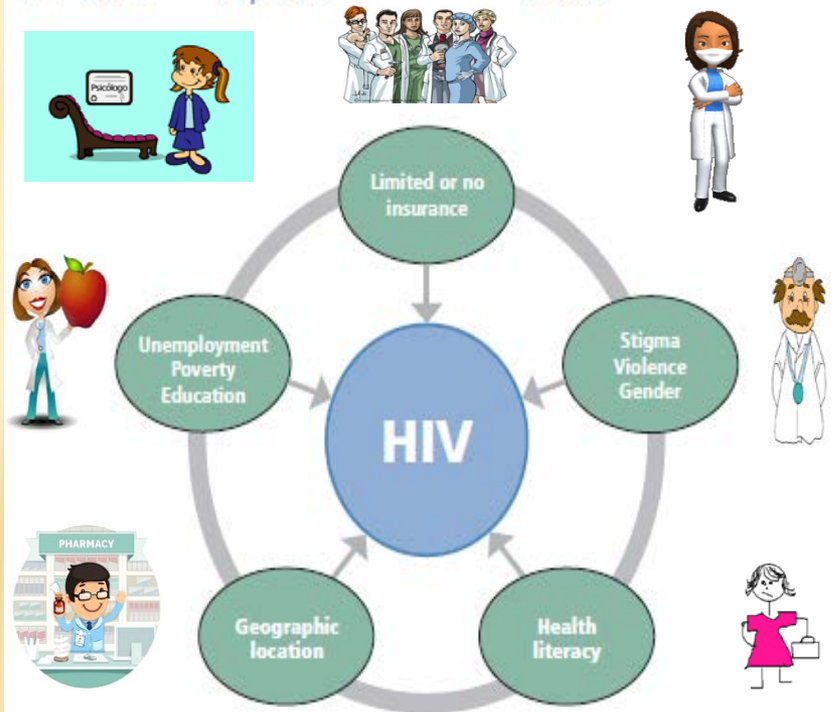
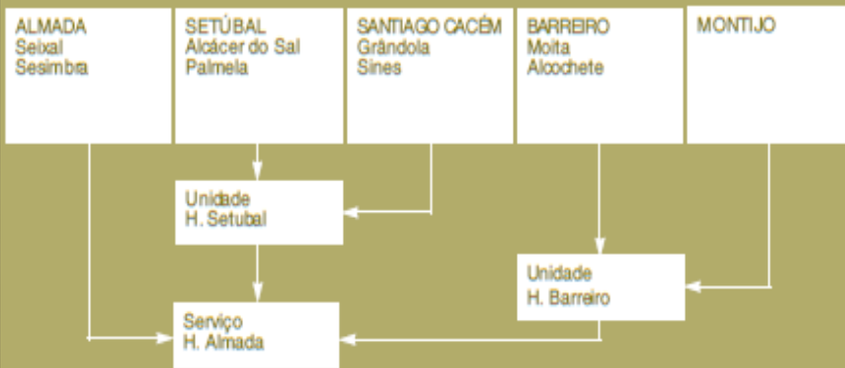


Figure 2. HIV infection: A single factor among many potential socioecologic disparities.

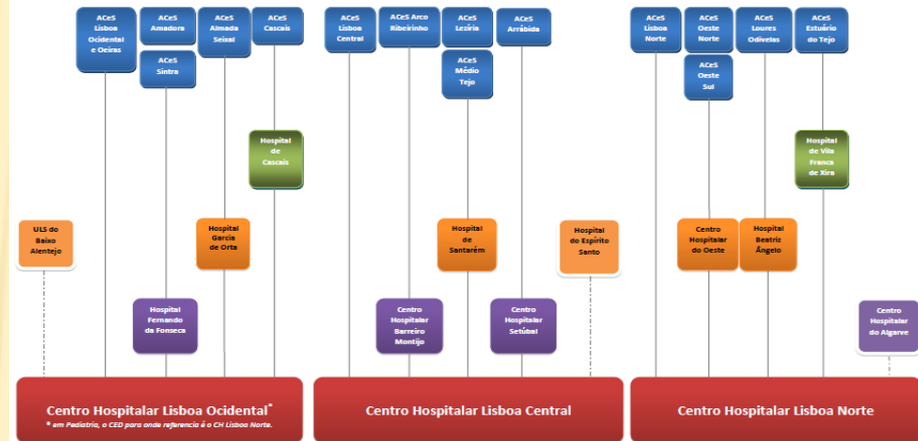
... ATRAVÉS DA DESCENTRALIZAÇÃO DE ALGUNS CUIDADOS DE AMBULATÓRIO...

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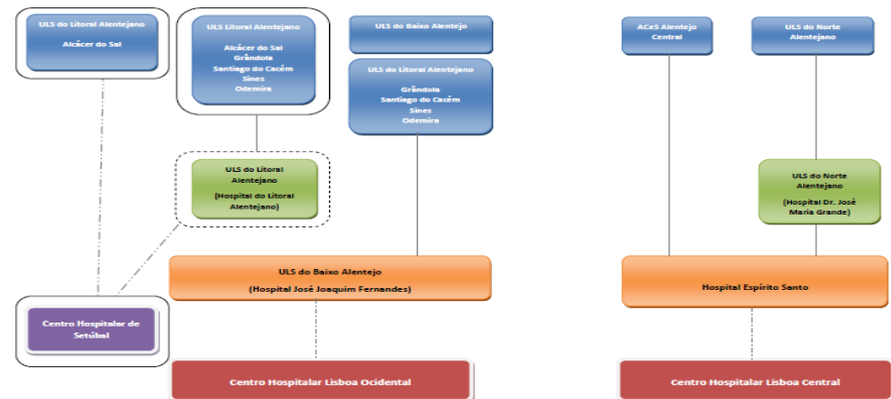
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Região de Lisboa e Vale do Tejo



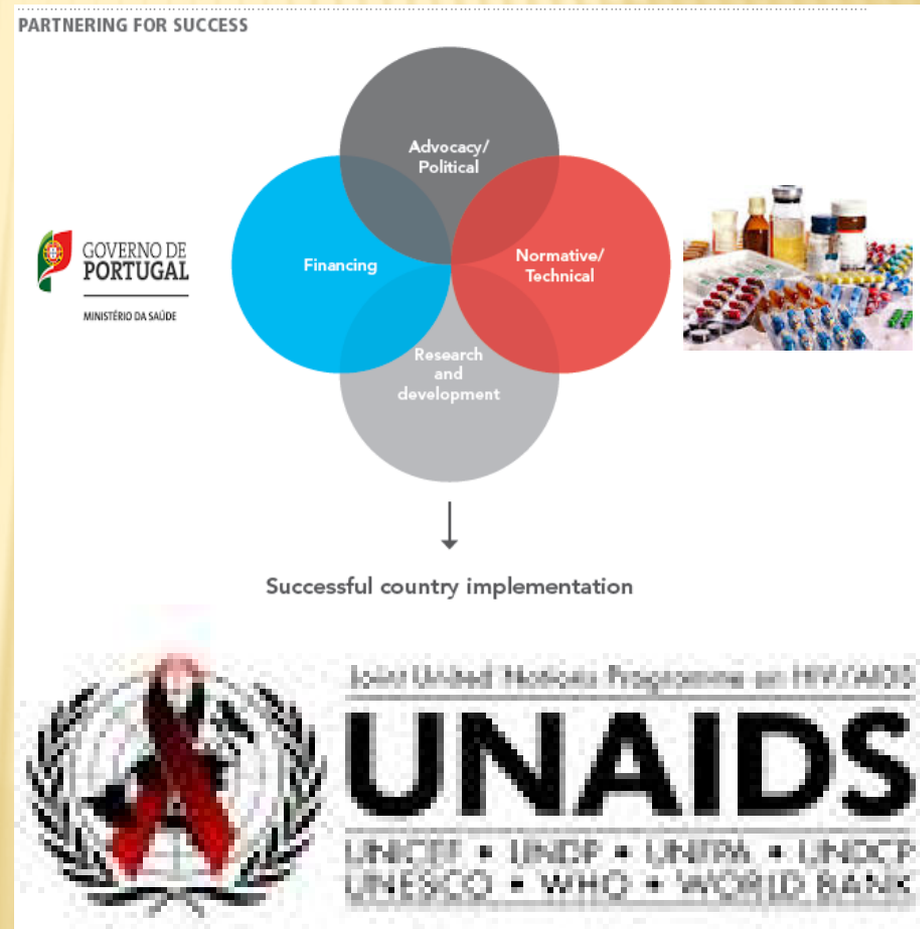
Região do Alentejo



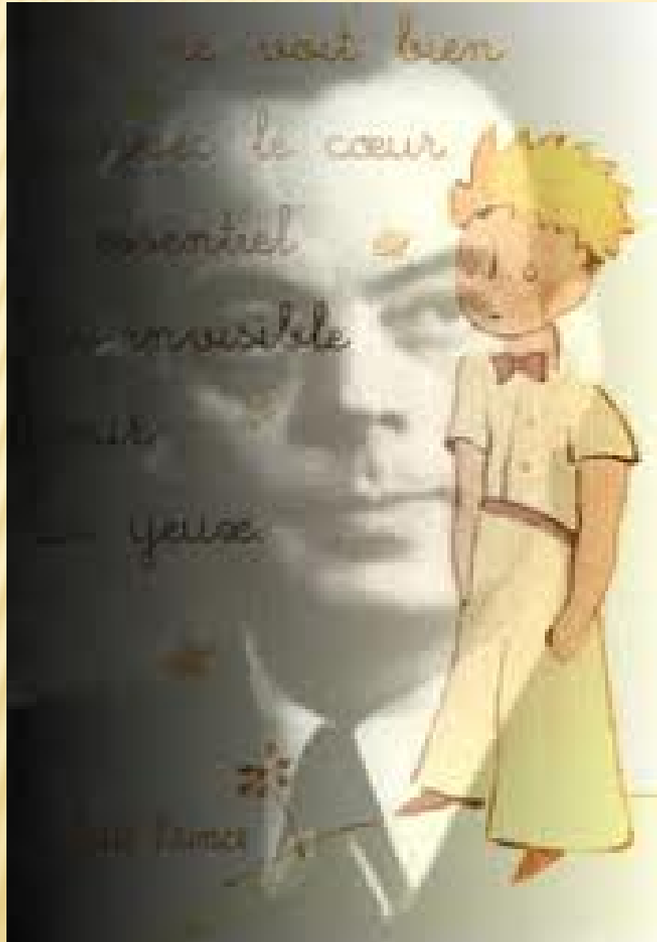
... (RE)AFIRMANDO QUE A SUSTENTABILIDADE PASSA CERTAMENTE POR IMPLEMENTAR ESTAS MEDIDAS!!!

✘ Propostas

- + 1)- Compra centralizada dos Medicamentos
- + 2)- Preço de referência igual p/ fármacos do mesmo grupo farmacológico e geração
- + 3)- Preço das co-formulações idêntico ao da soma dos seus componentes (incluindo genéricos) acrescido de um coeficiente justo (10%?) a ser negociado entre MS e IF
- + 4)- Estabelecimento de um nº máximo de doentes a serem tratados anualmente para as patologias que utilizem fármacos inovadores c/ impacto económico significativo (acima desse valor a IF suportaria os custos)
- + 5)- O pagamento ao Hospital prestador deverá acompanhar o doente (e a doença) permitindo assim a livre escolha por parte do doente sem mais constrangimentos
- + 7)- Informatização adequada do PC e avaliação periódica de resultados



UMA DERRADEIRA E INQUIETANTE INTERPELAÇÃO



- × *“Se a vida não tem preço, nós comportamo-nos sempre como se alguma coisa ultrapassasse, em valor, a vida humana... Mas o quê?”*

de Saint-Exupéry)

(Antoine

PARA QUE ESTAS IMAGENS NÃO TENHAM QUE VOLTAR A REFLETIR A REALIDADE



**TESE: O ESTADO SOCIAL SÓ NÃO ACABA PORQUE
VIVEMOS EM DEMOCRACIA E NENHUM POLÍTICO
GANHARIA ELEIÇÕES SE PROPUSESSE ABERTAMENTE A
SUA EXTINÇÃO**

