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**REPORTING
'LAB' RESULTS**

**THE CAUSE OF, AND
THE SOLUTION TO**

**THE OVERDIAGNOSIS
PROBLEM**

You can find a pdf of the handouts at
<http://therapeuticseducation.org/handouts>

Objectives

- 1) outline the problem of lab test measurement and reporting and some of the ways it contributes to the overdiagnosis problem
- 2) demonstrate with some examples (BP, LDL, glucose, bone density)
- 3) hopefully offer some useful tips, and suggestions and simple charts for how to deal with this extremely important and relevant healthcare conundrum
- 4) INTERACTIVE
 - Poll questions - internet access
 - Play with dice - work through a few scenarios

Just a few of the diagnoses that are solely or partially lab-based dependent

| | | | | | |
|---|---|--|--------------------------------------|--|--|
| Acid-Base Disorders | Celiac Sprue | Folic Acid or B9 Deficiency | Inhalation anthrax | Nephrotic Syndrome | SLE |
| Acidosis and Alkalosis | Cervical Cancer | Food and Waterborne Illness | Inherited Copper Toxicity | Neural Tube Defects | Small Cell Lung Cancer |
| Acidosis/Alkalosis | CF | Food Poisoning | Insulin Resistance | Neuropathy | Spina bifida |
| aCL Syndrome | CFIDS | Fungal Infections | Insulin Resistance Syndrome | NHL | Spinal dysraphism |
| ACS | CFS | Gastroenteritis | Iron Overload Disease | Non-Hodgkin lymphoma | Spinal Meningitis |
| Acute DIC | CHF | Gluten-Sensitive Enteropathy | Iron Storage Disease | Non-Small Cell Lung Cancer | SSc |
| Acute Idiopathic Polyneuritis | Chlamydia | Gonorrhea | Jaundice | Nontuberculous Mycobacteria | Stable angina |
| Acute Inflammatory Demyelinating | Chronic Fatigue and | Gout | JIA | Nontuberculous Mycobacteria Infections | Staph |
| Polyneuropathy | Immune Dysfunction Syndrome | Gouty Arthritis | JRA | NTD | Staph aureus |
| Acute Kidney Injury | Chronic Fatigue Syndrome | Graves Disease | Juvenile Idiopathic Arthritis | NTM | Staph Infections |
| Acute Myocardial Infarct | Chronic Kidney Disease | GSE | Juvenile Rheumatoid Arthritis | OA | Staph Infections and Methicillin-Resistant |
| Acute Renal Failure | Chronic Thyroiditis | Guillain-Barré Syndrome | Keratoconjunctivitis Sicca | Obesity Syndrome | Staphylococcus aureus |
| AD | Circumscribed Scleroderma | H1N1 | Kidney Disease | Osteoarthritis | Staphylococcus aureus |
| Addison Disease | Cirrhosis | H3N2 | Lactase Deficiency | Osteoarthritis | STDs |
| Adrenal Insufficiency | CKD | H5N1 | Lactose Intolerance | Osteoporosis | Stein-Leventhal Syndrome |
| Adrenal Insufficiency and Addison Disease | Coagulopathy | H7N9 | Landry's Ascending Paralysis | Ovarian Cancer | Sticky Blood Syndrome |
| AKI | Cobalamin Deficiency | Hashimoto Thyroiditis | LE | PA | STIs |
| Albuminuria | Colon Cancer | HBP | Lead Poisoning | Pancreatic Cancer | Stomach Flu |
| Alcohol dependence | Colorectal Cancer | HD | Leukemia | Pancreatic Diseases | Stroke |
| Alcoholism | Community-Acquired Pneumonia | Healthcare-Associated Pneumonia | Limited Cutaneous Scleroderma | Pancreatic Insufficiency | Subacute Cutaneous Lupus |
| Allergies | Congenital Adrenal Hyperplasia | Heart Attack | Linear Scleroderma | Pancreatitis | Swine Flu |
| Alzheimer Dementia | Congenital Alactasia | Heart Attack and Acute Coronary Syndrome | Liver Disease | Parathyroid Cancer | Syndrom X |
| Alzheimer Disease | Congestive Heart Failure | Heart Disease | Lobar Pneumonia | Parathyroid Diseases | Syphilis |
| AMI | Conn Syndrome | Heart Failure | Localized Scleroderma | PCOS | Systemic Exertion Intolerance Disease |
| Anemia | Consumption Coagulopathy | Hematuria | Lower Respiratory Tract Infection | Pelvic Inflammatory Disease | Systemic Lupus Erythematosus |
| Anencephaly | Copper Storage Disease | Hemochromatosis | Lung Cancer | Peptic Ulcer | Systemic Scleroderma |
| Angitis | CREST | Hemoglobin Abnormalities | Lung Diseases | PID | Systemic Sclerosis |
| Angina | Crohn Disease | Hemoglobin Barts | Lupus | Pituitary Disorders | TB |
| Angina pectoris | Cushing Syndrome | Hemoglobin C Disease | Lupus Anticoagulant Syndrome | Plasma Cell Dyscrasia | Testicular Cancer |
| Ankylosing Spondylitis | Cutaneous anthrax | Hemoglobin E Disease | Lupus Erythematosus | Plasma Cell Myeloma | Thalassemia |
| Anthrax | CVD | Hemoglobin S | Lyme Disease | Plasma Cell Neoplasm | Thrombophilia |
| Anticardiolipin Antibody Syndrome | Cystic Fibrosis | Hemoglobin Variants | Lymphocytic Thyroiditis | Plasmacytoma | Thyroid Cancer |
| Antiphospholipid Antibody Syndrome | Degenerative Joint Disease | Hemoglobinopathy | Lymphoma | Plasmacytoma of Bone | Thyroid Diseases |
| Antiphospholipid Syndrome | Dehydration | Hepatic Disease | Malabsorption | Pneumonia | Toxemia |
| aPL Syndrome | Dermatosclerosis | Hepatitis | Malaria | Polycystic Ovary Syndrome | Toxic Diffuse Goiter |
| APLS | Diabetes | Hepatolenticular Degeneration | Malignancy | Porphyria | Travelers' Diseases |
| APS | Diabetes mellitus | Hereditary Persistence of Fetal Hemoglobin | Malignant tumor | Post-infectious Arthritis | Trich |
| ARF | Diarrhea | Herpes | Malnutrition | Pre-eclampsia | Trichomonas |
| Arteritis | DIC | Herpes Zoster | MDS | Pregnancy | Trichomoniasis |
| Arthritis | Diffuse Cutaneous Scleroderma | High Blood Pressure | ME | Pregnancy-induced Hypertension | Trisomy 21 |
| AS | Diffuse Thyrotoxic Goiter | HIV | Melanoma | Presenile Dementia | Tuberculosis |
| Asthma | Disaccharidase Deficiency | HIV Infection and AIDS | Meningitis and Encephalitis | Primary Aldosteronism | Types of Liver Disease |
| Atypical Mycobacteria | Discoïd Lupus | HL | Meningococcal Meningitis | Primary Hyperaldosteronism | Ulcerative Colitis |
| Atypical Pneumonia | Disseminated Intravascular Coagulation | Hodgkin Disease | Menopause | Prinzmetal's angina | Unstable angina |
| Autoimmune Diseases | Disseminated Intravascular Coagulopathy | Hodgkin Lymphoma | Metabolic Syndrome | Prostate Cancer | Urinary Tract Infection |
| Autoimmune Thyroiditis | Disseminated Lupus Erythematosus | Hospital-Acquired Pneumonia | MG | Protein in urine | UTI |
| Avian Flu | DJD | HPFH | MI | Proteinuria | Vaginal Infection |
| Bacillus anthracis infection | Double Pneumonia | HPV | Morphea | RA | Vaginitis and Vaginosis |
| Bacterial Arthritis | Down Syndrome | Hughes Syndrome | MOTT | Reactive Arthritis | Vaginitis/Vaginosis |
| Bacterial Vaginosis | Drug-induced Lupus | Huntington Disease | MPDs | Reaven Syndrome | Variant angina |
| Benign Prostatic Hyperplasia | DS | Huntington's Chorea Disease | MPNs | Renal Disease, Kidney Failure | Vasculitis |
| Benign Prostatic Hypertrophy | Dysmetabolic Syndrome | Hypercoagulable Disorders or States | MRSA | Rheumatoid Arthritis | VD |
| Biological Warfare | Ebola Hemorrhagic Fever | Hyperparathyroidism | MS | Rheumatoid Spondylitis | Venereal Diseases |
| Bioterrorism Agents | Ebola Virus Disease | Hypersensitivity | Multiple Myeloma | Sarcoidosis | Vitamin B12 and Folate Deficiencies |
| Bleeding Disorders | Ebola Virus Infection | Hypertension | Multiple Sclerosis | SCD | Vitamin B12 Deficiency |
| Blood in the urine | Encephalitis | Hyperthyroidism | Myalgic Encephalomyelitis | Scleroderma | Vitamin K Deficiency |
| Bone Marrow Disorders | End Stage Renal Disease | Hypoparathyroidism | Myasthenia Gravis | SEID | Vulvovaginitis |
| Borrelia burgdorferi Infection | Endocrine Syndromes | Hypothyroidism | Mycobacteria other than tuberculosis | Seizure Disorder | Walking Pneumonia |
| Borrelia mayonii Infection | Endocrine System and Syndromes | IBD | Mycoses | Sepsis | West Nile Virus |
| BPH | Epilepsy | Icterus | Myelocoele | Septic Arthritis | Wilson Disease |
| Breast Cancer | ESRD | Infectious Arthritis | Myelodysplasia | Sexually Transmitted Diseases | WNV |
| CAH | EVD | Infectious Polyneuritis | Myelodysplastic Syndrome | Sexually Transmitted Infections | Wound and Skin Infections |
| Cancer | Excessive Clotting Disorders | Infertility | Myelomenigocele | Shingles | |
| Candidiasis | Extraosseous Plasmacytoma | Inflammatory Bowel Disease | Myeloproliferative Disorders | Sicca Syndrome | |
| Carbohydrate Intolerance | Fibromyalgia | Influenza | Myeloproliferative Neoplasms | Sickle Cell Anemia | |
| Cardiovascular Disease | Flu | Influenza A | Myocardial Infarct | Sickle Cell Disease | |
| Celiac Disease | Folate Deficiency | Influenza B | Neonatal Lupus | Sjögren Syndrome | |

“It is commonly thought that laboratory tests provide two-thirds to three-fourths of the information used for making medical decisions. If so, test results had better tell the truth about what is happening with our patients.”

Clinica Chimica Acta 2004;346:3-11

New Rule Grants Patients Direct Access to Lab Results

By Melinda Beck

Feb. 3, 2014 1:05 p.m. ET

Clinical laboratories must give patients access to their own lab-test results upon request, without going through the physician who ordered them, according to a new federal rule announced Monday by the Department of Health and Human Services.



PROBLEM #1

It's typically the same report that goes to health care providers

PROBLEM #2

Many health care providers don't appreciate the key nuances of "lab" tests

MY THESIS

“For much in medicine, we knowingly sell preeminent precision even though we all know in our heart of hearts we can only deliver educated estimates.

I believe most patients would be very understanding about this imprecision if we were just more open about it.”

-James McCormack, Pharm D (1959 - hopefully not soon)

“We also CAN’T be precise about the imprecision”

I am speaking in general, and do realise there are always some exceptions

I am presenting concepts

I will be providing ball-park estimates

Two Problems with Faking Precision



FALSE BELIEFS

BELIEF #1 - the good/bad thresholds are relatively black and white

BELIEF #2 -when the numbers change these changes are real

These beliefs can potentially lead to inappropriate feelings
of fear, happiness, frustration, confusion...

Both in patients AND clinicians

Sources of Imprecision

Lab
Error

Analytic
variation

Biologic
variation

Actual LAB errors

Lab
Error

0.3% 

- ~60% pre-analytical
- ~15% analytical
- ~25% post analytical

Table 1. Laboratory errors in stat testing.

| Defects: detection steps | Defects found | |
|--|---------------|--------------|
| | No. | Frequency, % |
| Preanalytical | | |
| Specimen collected from infusion route | 3 | 1.9 |
| Sample contaminated | 1 | 0.6 |
| Tube filling error | 21 | 13.1 |
| Empty tube | 11 | 6.9 |
| Inappropriate container | 13 | 8.1 |
| Nonrefrigerated sample | 3 | 1.9 |
| Missing tube | 5 | 3.1 |
| Digoxin test timing error | 1 | 0.6 |
| Patient identification error | 14 | 8.8 |
| Request procedure error | 12 | 7.5 |
| Data communication conflict | 6 | 3.8 |
| Physician's request order missed | 3 | 1.9 |
| Order misinterpreted | 2 | 1.3 |
| Check-in not performed (in the Laboratory Information Systems) | 4 | 2.5 |
| Subtotal | 99 | 61.9 |
| Analytical | | |
| Instrument-caused random error | 3 | 1.9 |
| Analytical inaccuracy not recognized | 21 | 13.1 |
| Subtotal | 24 | 15 |
| Postanalytical | | |
| Results communication breakdown | 32 | 20 |
| Lack of communication within laboratory | 3 | 1.9 |
| TAT excessive | 2 | 1.3 |
| Subtotal | 37 | 23.1 |

Clinical Chemistry 2007;53:1338-42

Dispensing errors ~1-2%

Measurement Landscape

Assuming no pre-analytic issues - timing/labelling etc

Population-based reference intervals

Analytical Variation

Analytic
variation

CVA - analytical variation

Biological Variation

Biologic
variation

CVI - within subject

CVG - between subject

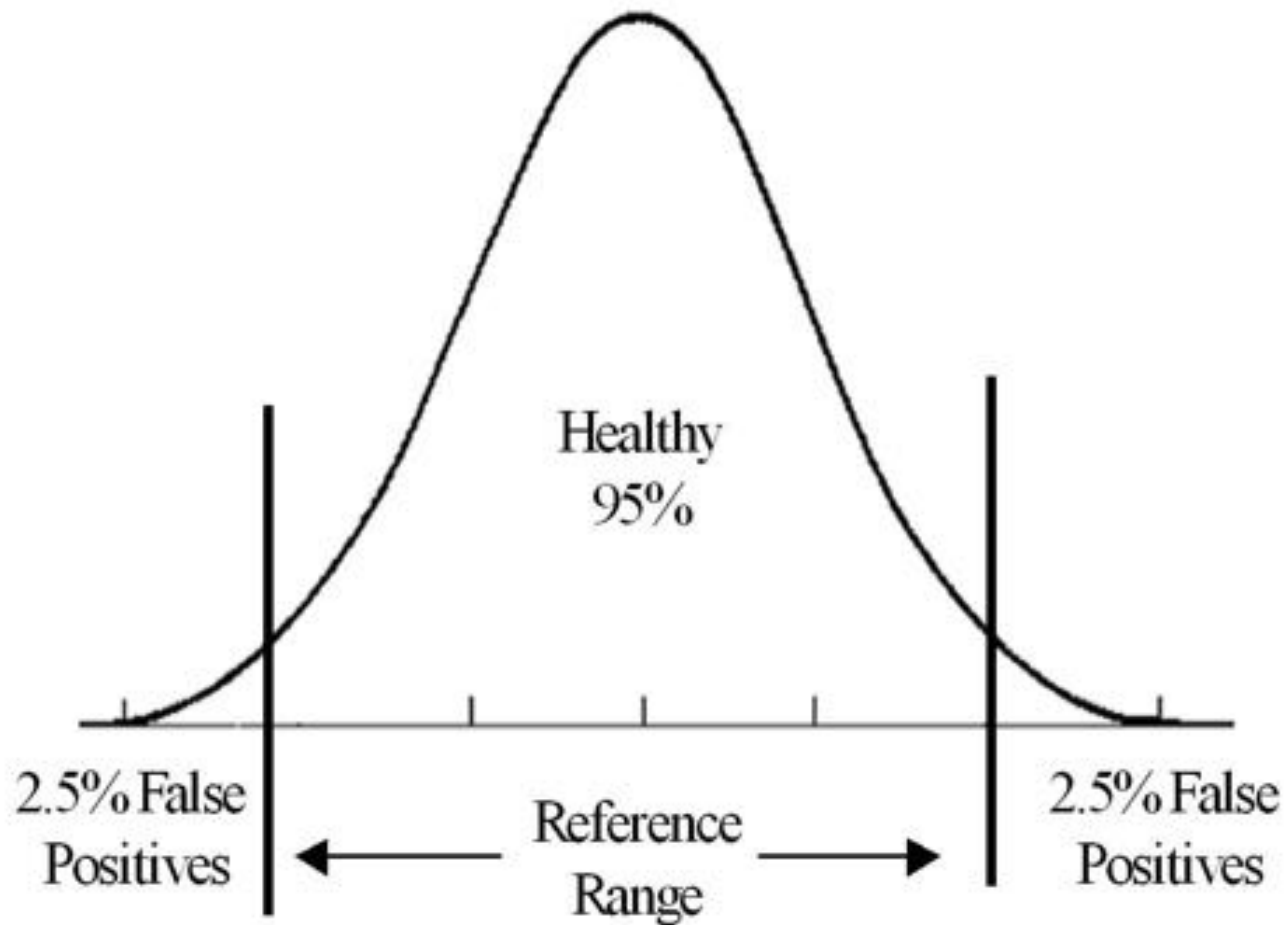


Reference change values (RCV)

Population-based
reference intervals

Population-based reference intervals

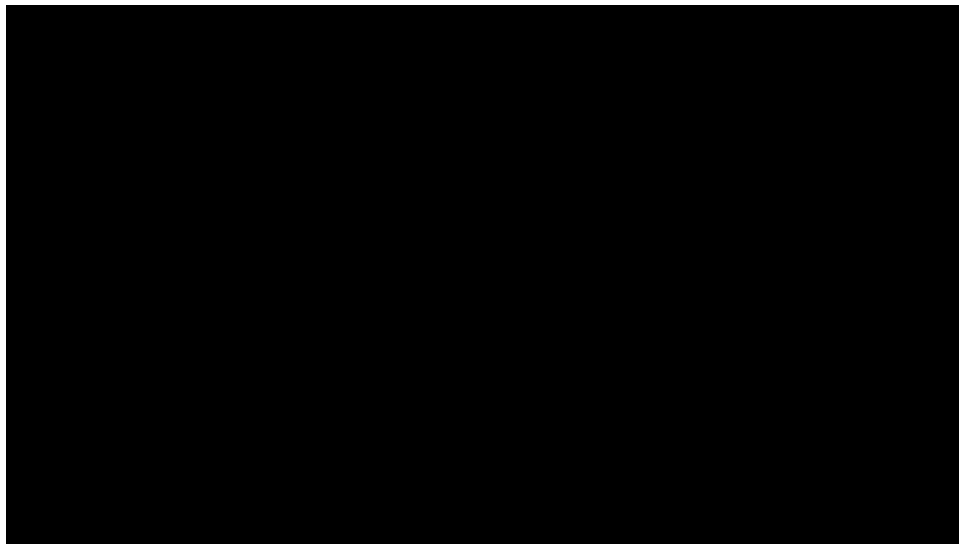
The interval/range where 95% of healthy people fall



Lab results report
exact numbers

BUT

Every test result is
really only a range
that hopefully includes
the true result
*+/- 1-2% up to
+/-20-30% or more*



YOU CANNOT BE SERIOUS!!
That ball was on the line

| Number of Tests Ordered | Probability of at Least One Abnormal Test |
|-------------------------|---|
| 1 | 5% |
| 2 | 10% |
| 5 | 23% |
| 10 | 40% |
| 15 | 54% |
| 20 | 64% |

When we do tests, typically
we are wondering

what are the results NOW, and/or

have they changed from PREVIOUS
measurements

Analytic
variation

Biologic
variation

Every “measurement”
will be “different”

Analytic variability
Biologic variability



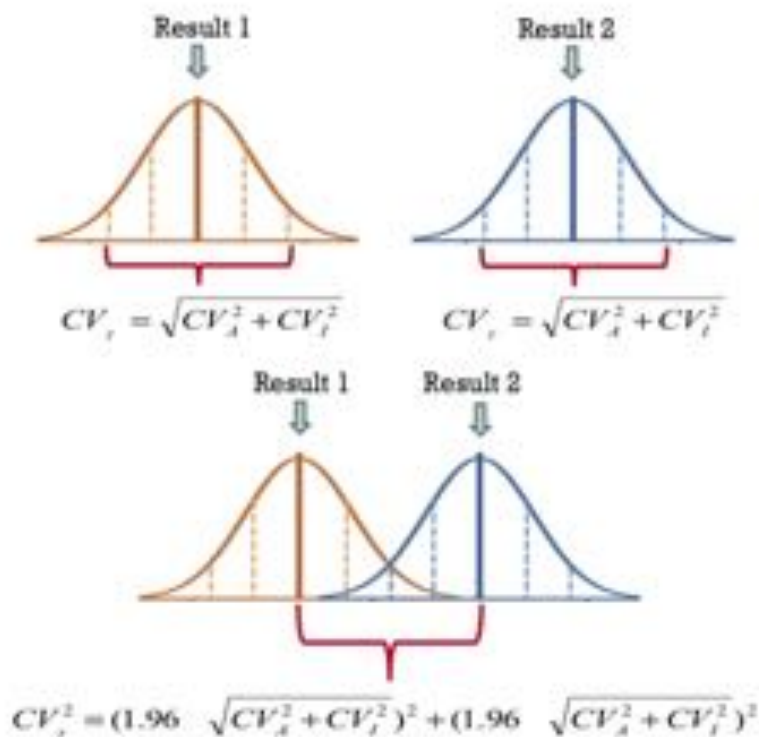
Reference Change Values (RCV)

a tool for assessment of the significance of differences
in serial results from an individual

Reference Change Values

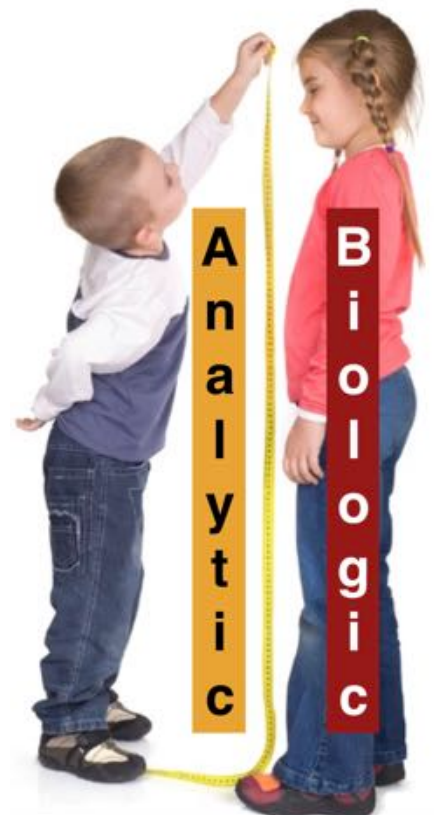
Used with SERIAL results to help deal with the analytic imprecision and biologic variation

Coefficients of Variation (total) = analytic PLUS biologic variation



MINIMUM DIFFERENCE
between two consecutive
results which needs to be
EXCEEDED in order for
one to state a
STATISTICALLY
SIGNIFICANT
change has taken place

$$RCV = \sqrt{2} * 1.96 * \sqrt{(CV_{Analytical}^2 + CV_{Intraindividual}^2)}$$



How good, analytically speaking,
does a “test” need to be

“The analytical CV (CVA) should be
less than one-half the average within-
subject biological variation (CVI)”



When it is, the CVA has almost
no impact on the RCV - the
RCV is pretty much determined
by the CVI



Reference change values provide a “p-value”
for the differences between two measurements

THE AMERICAN STATISTICIAN
2016, VOL. 70, NO. 2, 129–133
<http://dx.doi.org/10.1080/00031305.2016.1154108>



EDITORIAL

The ASA's Statement on p -Values: Context, Process, and Purpose

In February 2014, George Cobb, Professor Emeritus of Mathematics and Statistics at Mount Holyoke College, posed these (American Statistical Association 2010). However, these were

“It’s science’s dirtiest secret: The ‘scientific method’ of testing hypotheses by statistical analysis stands on a flimsy foundation.”

“Numerous deep flaws in null hypothesis significance testing.”

“Statistical techniques for testing hypotheses ...have more flaws than Facebook’s privacy policies.”

Experts issue warning on problems with P values

Misunderstandings about common statistical test damage science and society

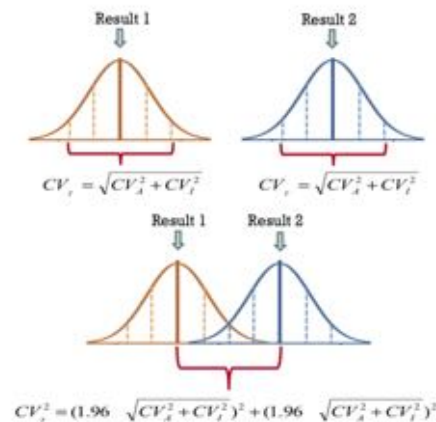
BY TOM SIEGFRIED 10:30AM, MARCH 11, 2016

Reference Change Values

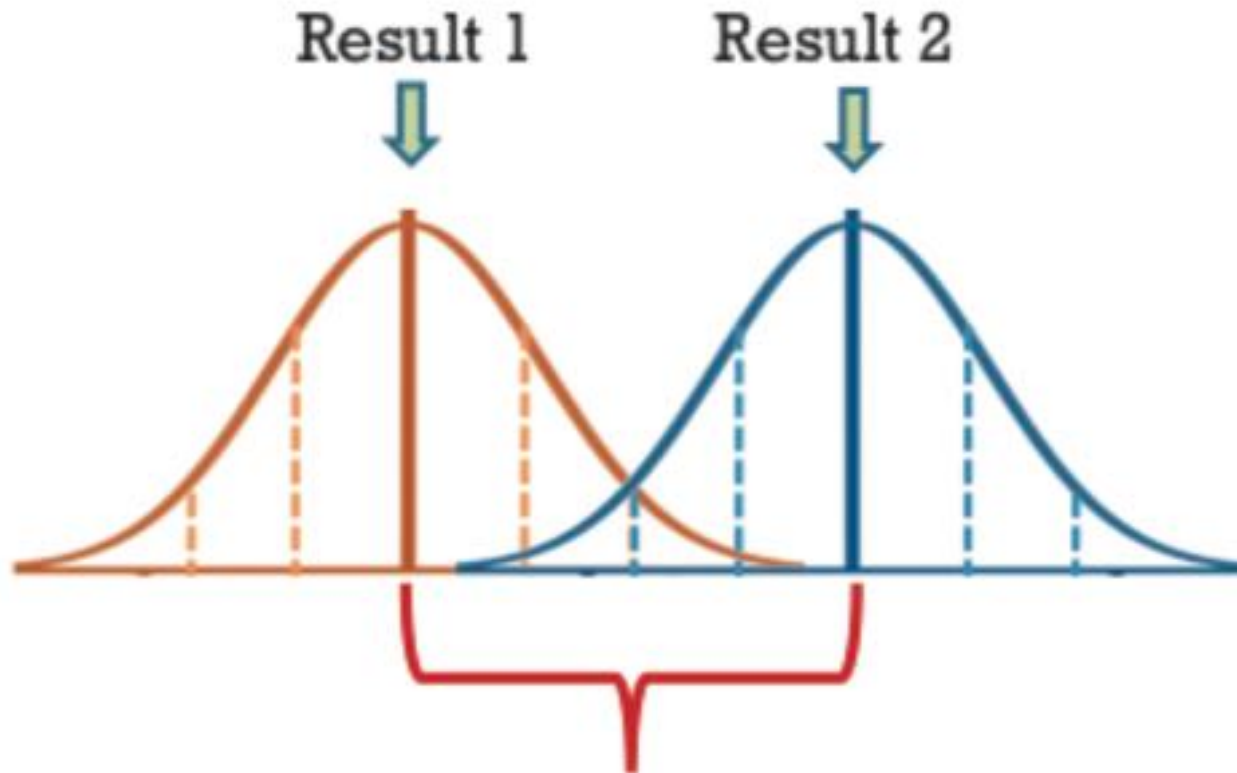
findings of a “significant difference” JUST means we are ruling out that the difference seen is due to chance

NOT

THAT THE MAGNITUDE OF THE DIFFERENCE SEEN IS THE ACTUAL MAGNITUDE OF THE DIFFERENCE



We believe these two results are different



can't necessarily quantify this
difference with any precision

What about multiple measurements?

Table 1. RCV using multiple estimates of the initial and new set points, expressed as a fraction of traditional RCV from two singleton measurements.

| | | Number of results estimating initial set point | | | | |
|---|---|--|------|------|------|------|
| | | 1 | 2 | 3 | 4 | 5 |
| Number of results estimating new set point | 1 | 1.00 | 0.87 | 0.82 | 0.79 | 0.77 |
| | 2 | 0.87 | 0.71 | 0.65 | 0.61 | 0.59 |
| | 3 | 0.82 | 0.65 | 0.58 | 0.54 | 0.52 |
| | 4 | 0.79 | 0.61 | 0.54 | 0.50 | 0.47 |
| | 5 | 0.77 | 0.59 | 0.52 | 0.47 | 0.45 |

with 4 measurements before and 4 afterwards
(vs 1 before and 1 after)
you can lower the RCV by 50%

Lab
Error

Analytic
variation

Biologic
variation



This is the
problem
and it is
NOT fixable,
it is only
KNOWABLE

Glucose
Blood pressure
Cholesterol
Bone Density





Glucose

Precisely Imprecise

What an A1c result really means

4.5% 5.8% 6.3% 6.8% 9%

| | | |
|-------------------------------|------------------------------|--|
| <p>Normal</p> <p>5</p> | <p>Pre-diabetes</p> <p>6</p> | <p>Diabetes</p> <p>7 8 9 10</p> |
|-------------------------------|------------------------------|--|

0.7%

A1c %

Typical A1c change seen
with a medication
= 0.7% ↓

Seasonal variation 0.2-0.5%
Higher in the winter

Am J Epi 2004;161:565-74

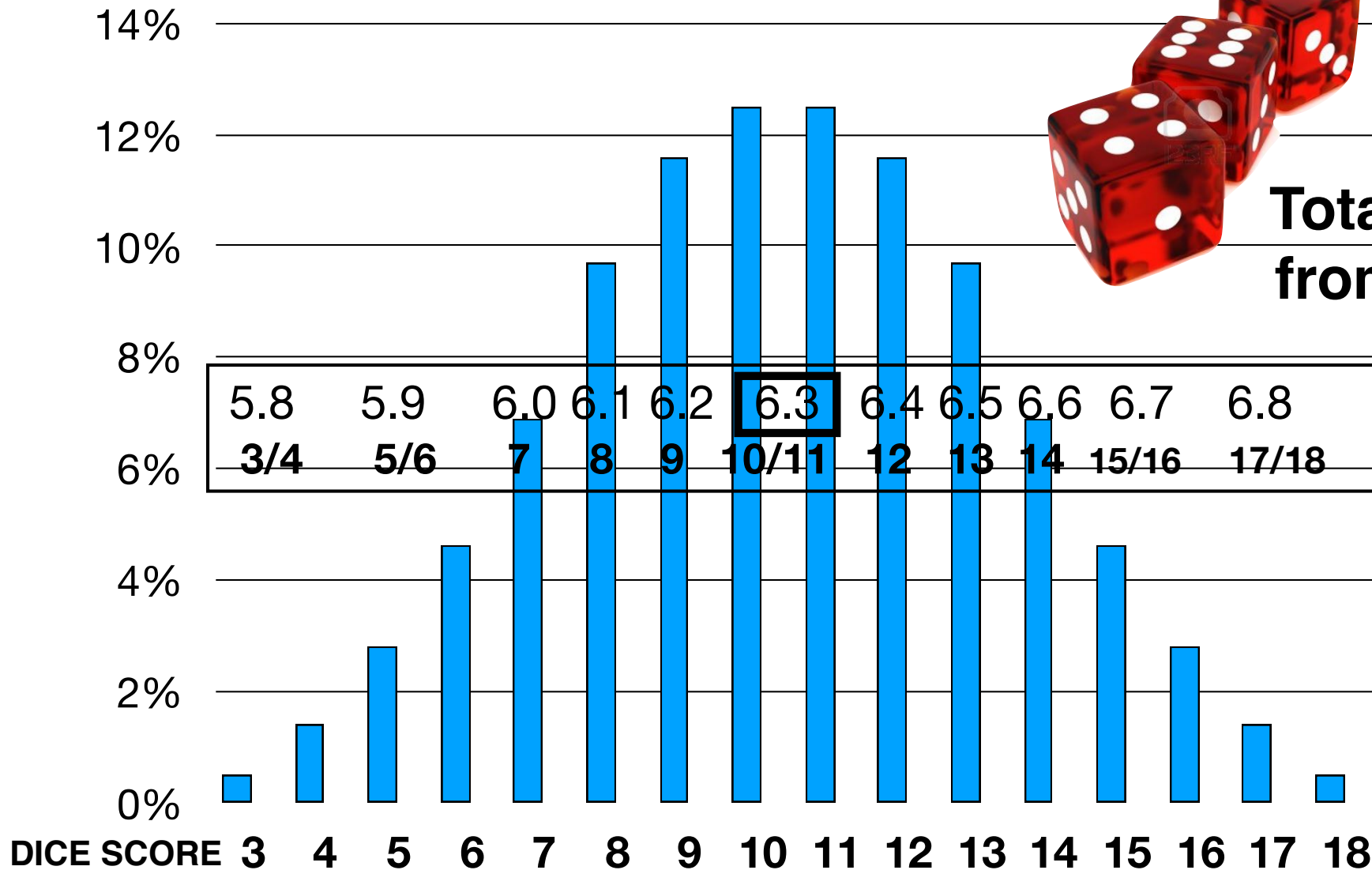
Assume your A1c
is definitely 6.3%

A1c %

= True value



Total score from 3 dice

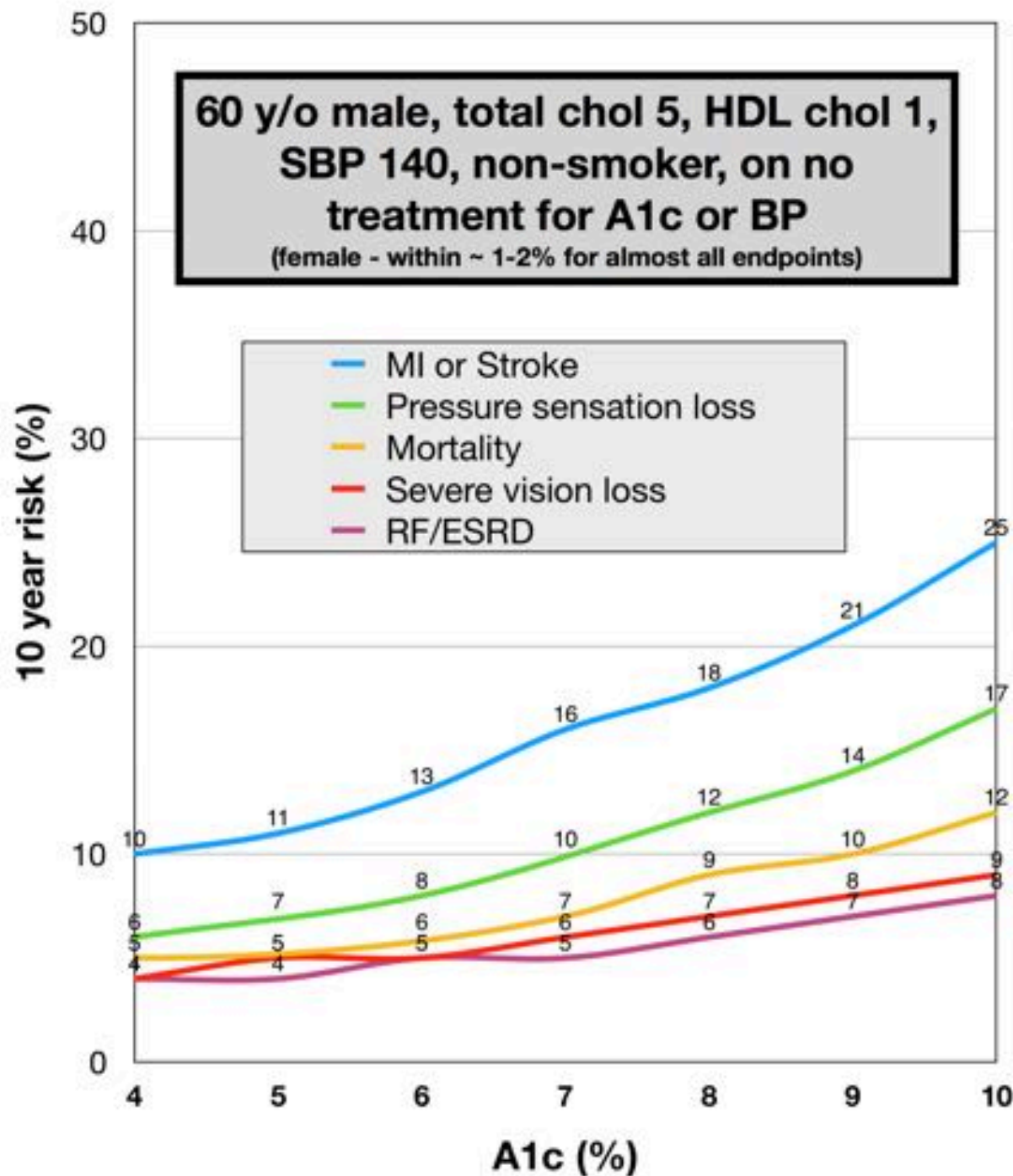


%

Normal
≤6.0

Pre-Diabetes
6.1-6.4

Diabetes
≥6.5



T2DM risk
should not
be
categorized
as
YES
or
NO



Blood pressure

Systolic blood pressure

TYPICAL CHANGES SEEN

Start medication - avg 9 mmHg ↓

Increase dose - avg 2-5 mmHg ↓

Seasonal differences - avg 8 mmHg ↓ when warm

Age related (per year) - avg 0.5-0.8 mmHg ↑

Sample size calculation - 40 office measurements before and after treatment to be REASONABLY confident that a 5 mmHg change has occurred



Editorial

American Journal of Hypertension (2008) 21:3–4; doi:10.1038/ajh.2007.20

Blood Pressure Variability: The Challenge of Variation

Tom P Marshall [1,✉](#)

¹Department of Public Health and Epidemiology, University of Birmingham, Edgbaston, Birmingham, UK

Correspondence: Tom P. Marshall, (T.P.Marshall@bham.ac.uk (mailto:T.P.Marshall@bham.ac.uk))

Need changes of at least 10/5 mmHg before you can say there has been a change

Am J Hyper 2008;21:3–4

“clinicians cannot identify individuals who have good or poor responses to drugs”

“coefficients of variation for systolic office, ambulatory, and self-monitored blood pressure, compared at baseline and 6 weeks, were 8.6%, 5.5%, and 4.2% respectively”

Br J Gen Pract 2010; 60: 675–80

“a single careful blood pressure measurement taken a few months after the start of treatment is not useful for monitoring”

BMJ 2009;338:b1492



Cholesterol

2016 Canadian Cardiovascular Society Guidelines for the Management of Dyslipidemia for the Prevention of Cardiovascular Disease in the Adult

“In individuals with a modified FRS of 5%-9%, yearly monitoring could be used to evaluate change in risk”

AACE 2017 Guidelines

AMERICAN ASSOCIATION OF CLINICAL ENDOCRINOLOGISTS AND AMERICAN COLLEGE OF ENDOCRINOLOGY GUIDELINES FOR MANAGEMENT OF DYSLIPIDEMIA AND PREVENTION OF CARDIOVASCULAR DISEASE

“Lipid status should be re-assessed 6 weeks after therapy initiation and again at 6-week intervals until the treatment goal is achieved.”

“While on stable lipid therapy, individuals should be tested at 6-to 12-month intervals”

Monitoring Cholesterol Levels: Measurement Error or True Change?

Paul P. Glasziou, MBBS, PhD; Les Irwig, MBBS, PhD; Stephane Heritier, PhD; R. John Simes, MBBS, MD; and Andrew Tonkin, MBBS, MD, for the LIPID Study Investigators

Background: Cholesterol level monitoring is a common clinical activity, but the optimal monitoring interval is unknown and practice varies.

Objective: To estimate, in patients receiving cholesterol-lowering medication, the variation in initial response to treatment, the long-term drift from initial response, and the detectability of long-term changes in on-treatment cholesterol level ("signal") given short-term, within-person variation ("noise").

Design: Analysis of cholesterol measurement data in the LIPID

of variation, 7%) to 0.60 mmol/L (23 mg/dL) (coefficient of variation, 11%), but it took almost 4 years for the long-term variation to exceed the short-term variation. This slow increase in variation and the modest increase in mean cholesterol level, about 2% per year, suggest that most of the variation in the study is due to short-term biological and analytic variability. Our calculations suggest that, for patients with levels that are 0.5 mmol/L or more (≥ 19 mg/dL) under target, monitoring is likely to detect many more false-positive results than true-positive results for at least the first 3 years after treatment has commenced.

Ann Intern Med 2008;148:656-61

VARIATION

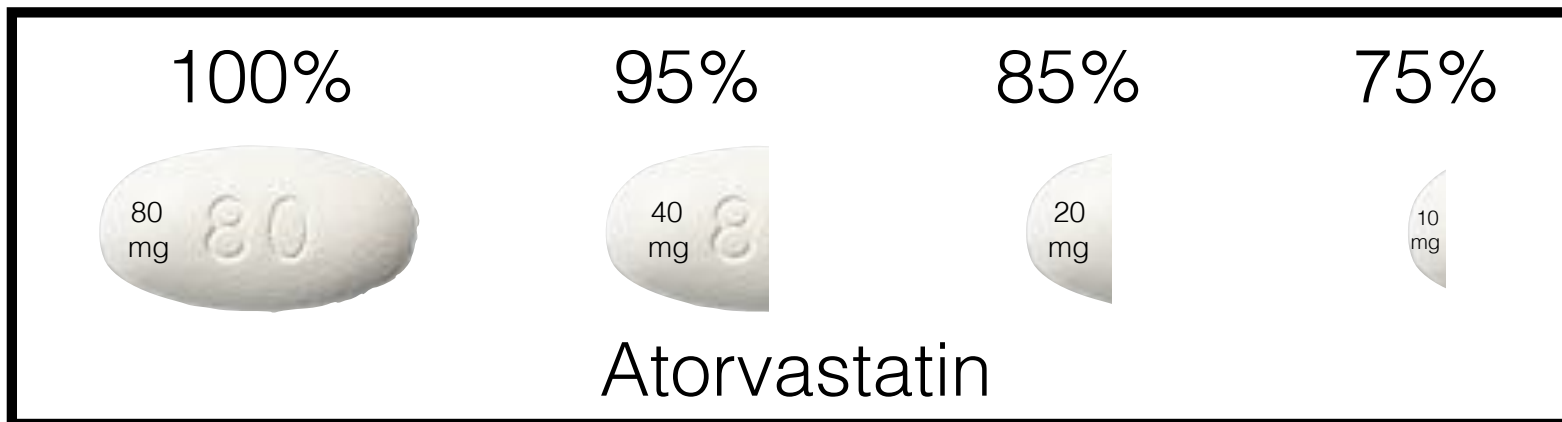
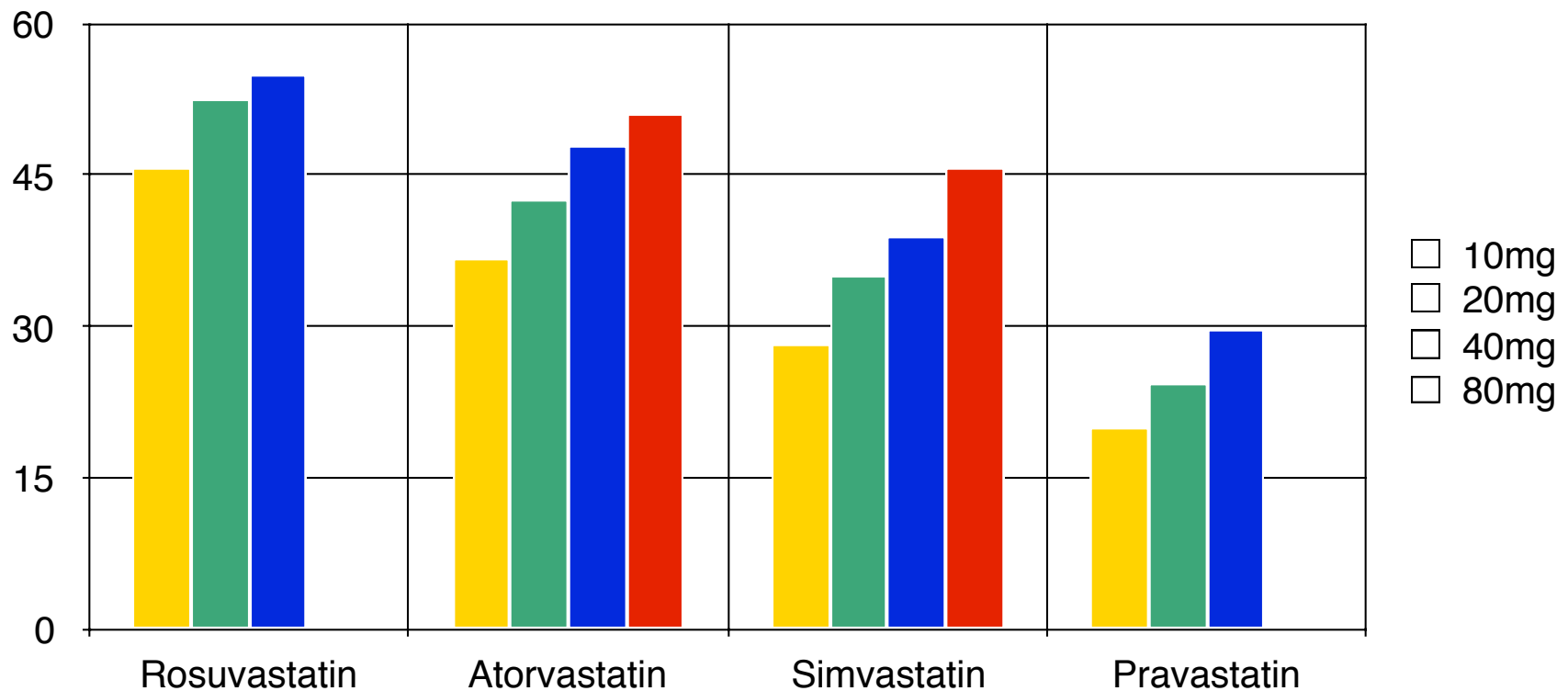
Total chol ~ - 0.80 to 0.80 mmol/L (~30 mg/dL)

LDL chol ~ - 0.5 to 0.5 mmol/L (~20 mg/dL)

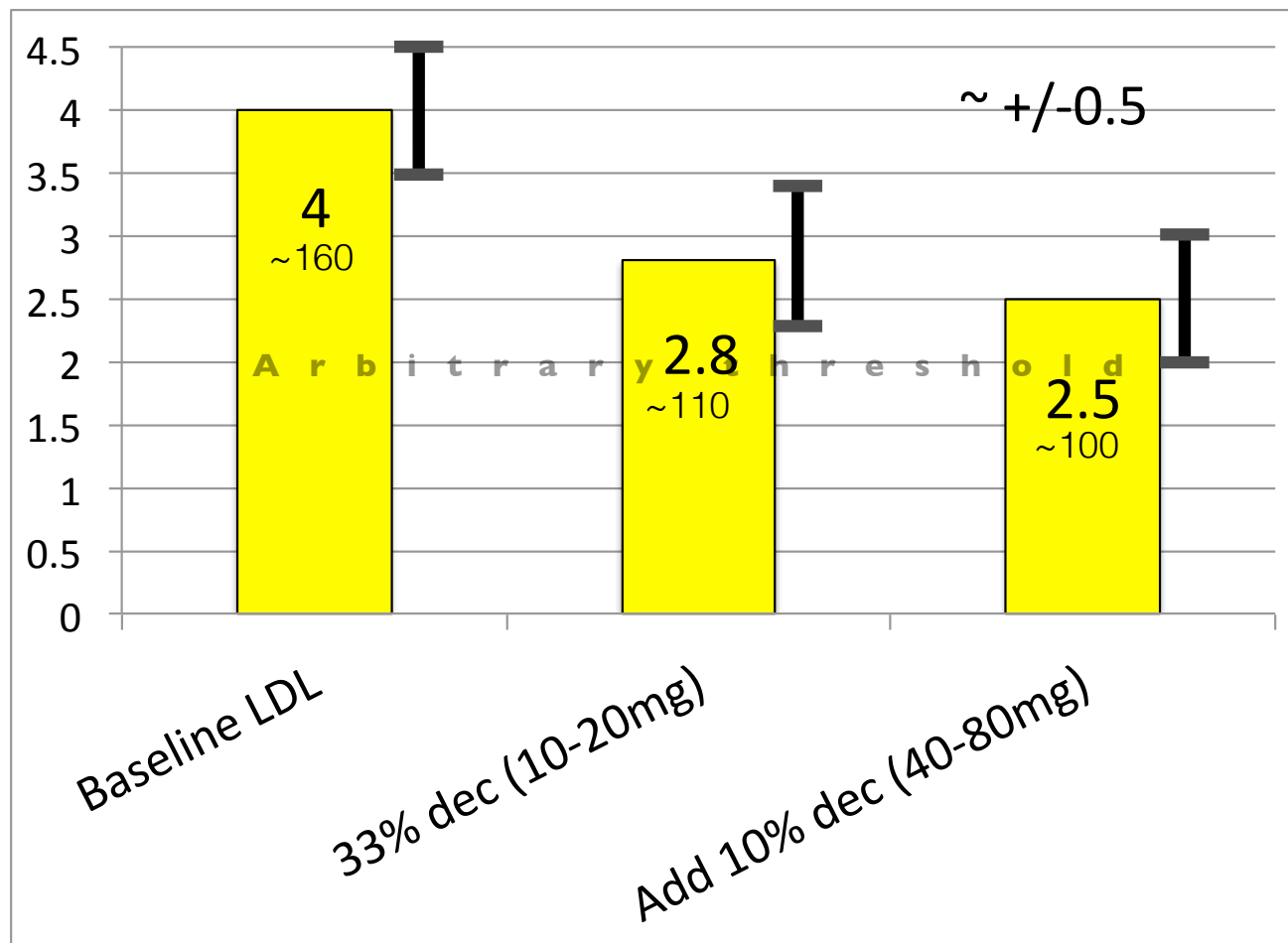
Average increase in cholesterol is 0.5-1%/year

“After initial change only measure every
3-5 years”

DOSE increases do not lead
to proportional EFFECT increases
% reduction in LDL cholesterol

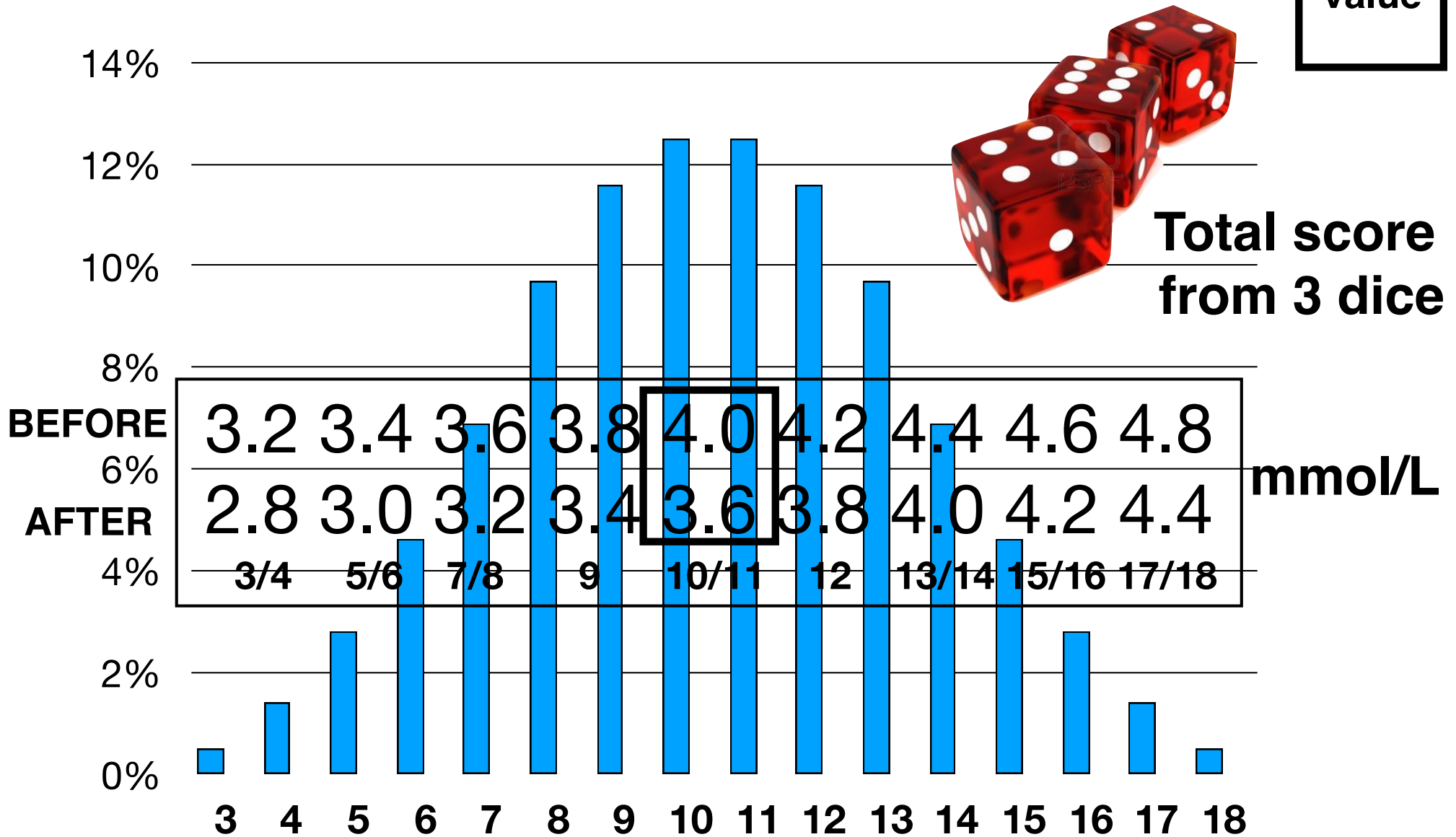


LDL cholesterol - 2 mmol/L ~80mg/dL



LDL mmol/L - medication that lowers LDL by 10%

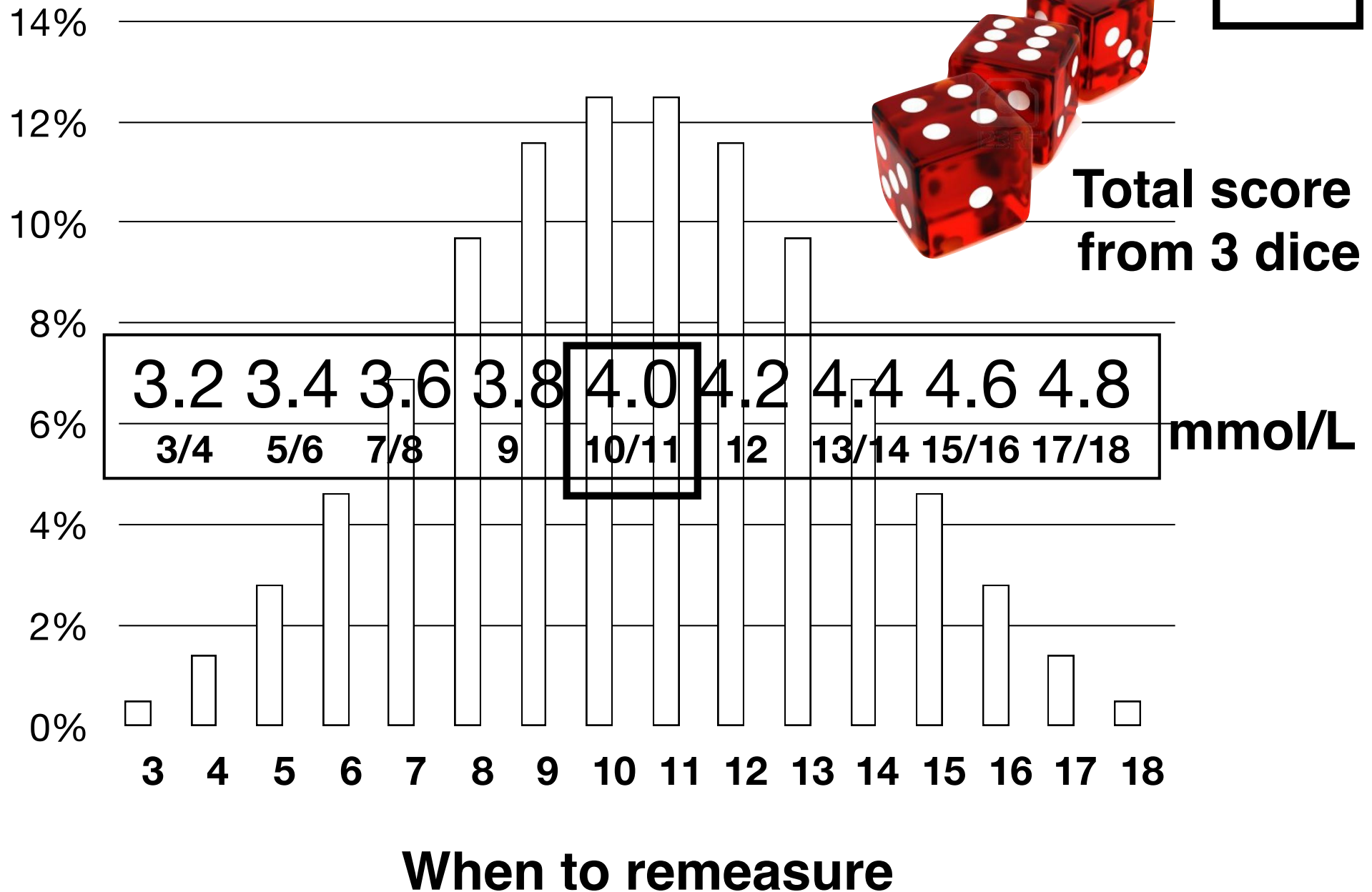
= True value



Trying to get to “target” by increasing the dose

LDL mmol/L - Average increase in cholesterol is 0.5-1%/year

**= True
value**



RESEARCH

When to remeasure cardiovascular risk in untreated people at low and intermediate risk: observational study

BMJ 2013; 346 doi: <http://dx.doi.org/10.1136/bmj.f1895> (Published 3 April 2013)

Cite this as: *BMJ* 2013;346:f1895

“Repeat risk estimation before 8-10 years is not warranted for most people initially not requiring treatment”

Languages: English (EN)

The Absolute CVD Risk/Benefit Calculator

Framingham
 US Data, 10 Year Risk
 Heart attacks + angina/coronary insufficiency + heart failure + strokes + intermittent claudication

QRISK®2-2014
 UK Data, 10 Year Risk
 Heart attacks + strokes

ACC/AHA ASCVD
 US Data, 10 Year Risk
 CHD death + nonfatal heart attacks + fatal/nonfatal strokes

PREDICT
 New Zealand Data, 5 Year Risk
 Heart attacks + angina + heart failure + strokes/TIAs + peripheral vascular disease

Age
 ——— 50 ——— years

Gender
 Male ☒ Female

Smoker
 Yes ☐ No ☒
CVD risk is reversed after 5-10 years of no smoking

Diabetes
 Yes ☐ No ☒

Systolic Blood Pressure
 ——— 120 ——— mmHg
Enter present blood pressure regardless of treatment
120 mmHg is used for baseline risk

On treatment for BP
 Yes ☐ No ☒
Click YES if taking blood pressure medication
Only applies if SBP is greater than 120 mmHg

Total Cholesterol
 ——— 3 ——— mmol/L
Cholesterol should be prior to drug treatment
3 mmol/L is used for baseline risk.
[Click to change to mg/dL.](#)

HDL Cholesterol
 ——— 1.3 ——— mmol/L
HDL should be prior to drug treatment
1.3 mmol/L is used for baseline risk.

Chronic Kidney Disease
 CKD status is not part of the risk algorithm but is used for calculating the benefit of certain therapies
 Yes ☐ No ☒

Relative Benefit: 0%
Benefit often has nothing to do with the effect on the surrogate marker. At present, you can only select one intervention at a time.

Physical Activity
 Mediterranean Diet vs Low fat
 Vitamin/Omega-3 supplements
 BP meds (not atenolol/doxazosin)
 Low-mid intensity statins
 High intensity statins
 Fibrates
 Niacin
 Ezetimibe
 Metformin
 Sulfonylureas
 Insulins
 Glitazones
 GLPs
 DPP-4s
 Meglitinides
 SGLT2
 Smoking Cessation
 ASA

[Benefit Estimate Details](#)

Risk Time Period
 10 years

😊 97.9% No event
 😞 2.1% Total with an event
 😊 0.0% Number who benefit from treatment
 NNT ∞ Number needed to treat
 😞 2.1% Baseline events using baseline factors alone
 😞 0.0% Additional events "caused" by risk factors

As with all risk calculators, calculated risk numbers are +/- 5% at best. [More information.](#)

Print Report

Calculate ballpark 5/10-yr risk of CVD - BP, chol, diabetes

Make estimate of benefit based on the best available evidence

Gives a list of adverse effects to discuss

cvdcalculator.com

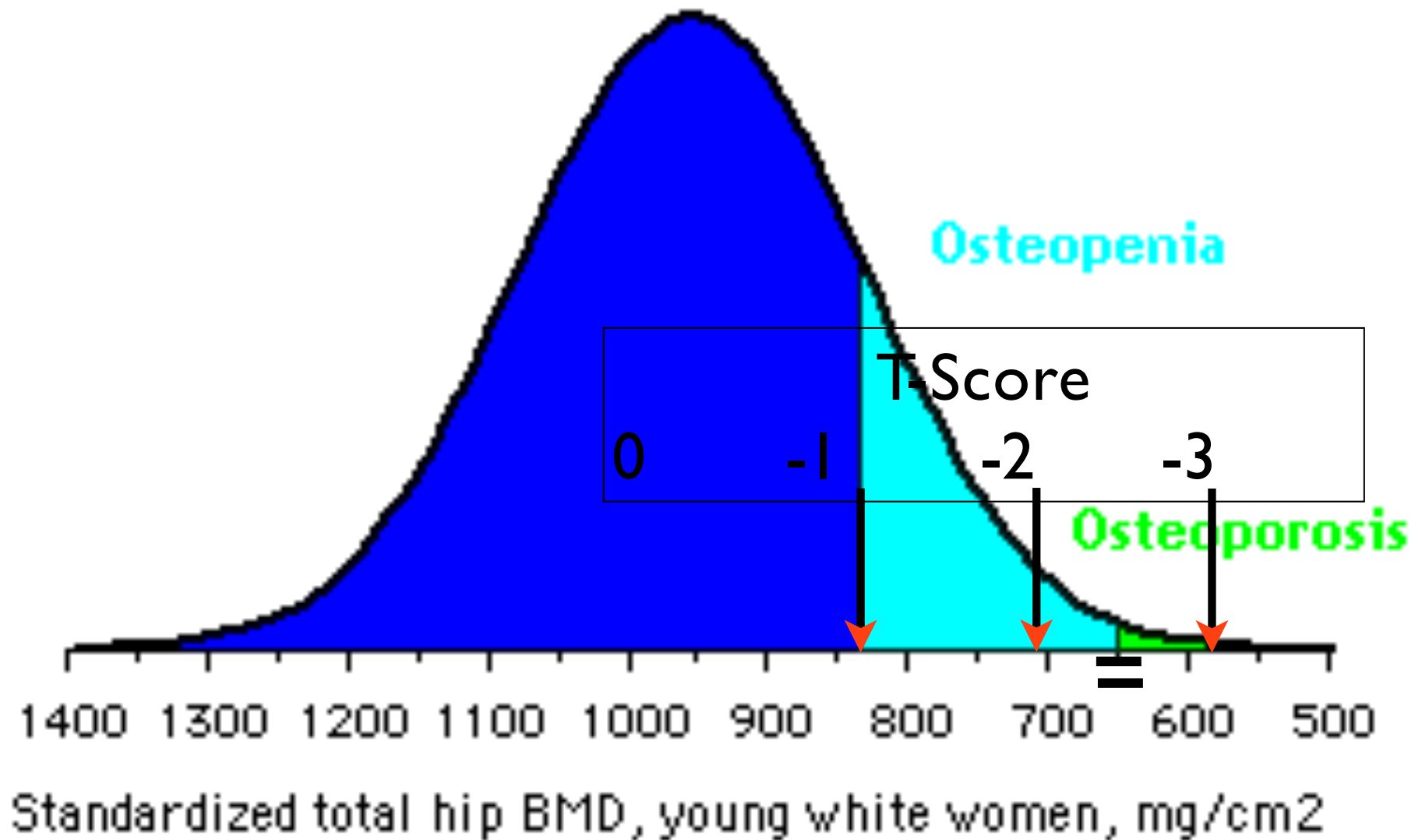


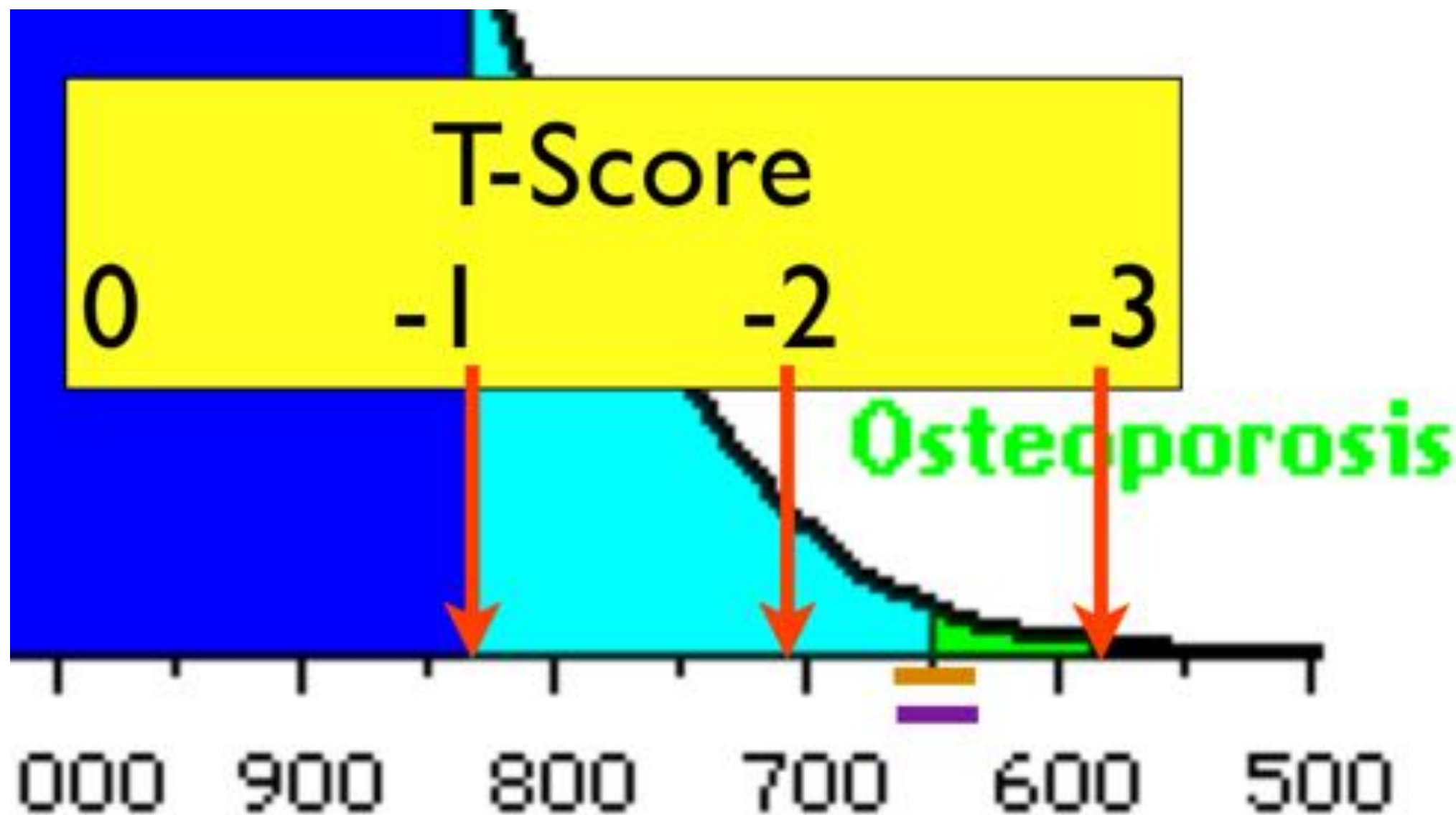
Bone density
(almost all analytic issues)

**AMERICAN ASSOCIATION OF CLINICAL ENDOCRINOLOGISTS
MEDICAL GUIDELINES FOR CLINICAL PRACTICE FOR
THE DIAGNOSIS AND TREATMENT OF
POSTMENOPAUSAL OSTEOPOROSIS 2010**

“Obtain a baseline DXA, and repeat DXA every 1 to 2 years until findings are stable. Continue with follow-up DXA every 2 years or at a less frequent interval”

- 1) Average bone loss per year $\sim 0.6\%$
- 2) Difference in BMD between drug and placebo - 3 years $\sim 5\%$
- 3) BMD measurement precision $\pm 2-3\%$





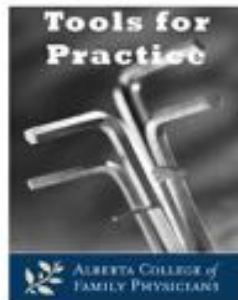
Other Smarter People

Value of routine monitoring of bone mineral density after starting bisphosphonate treatment: secondary analysis of trial data

Katy J L Bell, Andrew Hayen, Petra Macaskill, Les Irwig, Jonathan C Craig, Kristine Ensrud and Douglas C Bauer

BMJ 2009;338;b2266;

“Monitoring BMD in the first 3 years after starting treatment with a bisphosphonate is unnecessary and may be misleading”



Bone Mineral Density – Too much of a good thing?

Clinical Question: Once we have initiated bisphosphonate therapy, how frequently should we check bone mineral density (BMD)?

#32 Christina Korownyk & Michael R. Kolber

“Repeating BMD in the first three years after starting treatment with a bisphosphonate is unnecessary and potentially confusing. The vast majority of patients taking a bisphosphonate will get an adequate increase in BMD after three years and have a reduced fracture risk regardless of BMD changes”

Treatment of Low Bone Density or Osteoporosis to Prevent Fractures in Men and Women: A Clinical Practice Guideline Update from the American College of Physicians

“The data do not support monitoring BMD during the initial 5 years of treatment in patients receiving pharmacologic agents to treat osteoporosis.”

Other Smarter People

Average bone loss per year ~ 0.6%

Evaluating the Value of Repeat Bone Mineral Density Measurement and Prediction of Fractures in Older Women

The Study of Osteoporotic Fractures

Teresa A. Hillier, MD, MS; Katie L. Stone, PhD; Doug C. Bauer, MD;
Joanne H. Rizzo, MS; Kathryn L. Pedula, MS; Jane A. Cauley, DrPH;
Kristine E. Ensrud, MD, MPH; Marc C. Hochberg, MD; Steve R. Cummings, MD

Arch Intern Med. 2007;167(2):155-160.

“repeat BMD [8 years] measurement provides little additional benefit as a screening tool”

Arch Intern Med 2007;167:155-60

Now What?!!



“The obscure we see eventually.
The completely obvious, it seems,
takes longer.”

Edward R Murrow 1908-1965



The Problem
is
NOT Fixable,
it is
Only
KNOWABLE

Bottom Line



Embrace our “nudity”

Magnitude of the Imprecision Around Routinely Ordered Medical Measurements*

| | | | | | | |
|---|----------------------------------|--|---|--|--------------------------------|---|
| | Chloride Sodium Osmolality | Calcium Protein Bone density Hemoglobin A1 C Albumin Systolic BP | Magnesium Glucose Potassium pCO2 Cholesterol Creatinine Alk phos PTT | LDL HDL INR Total Cholesterol Phosphate LDH Uric acid Rheumatoid factor Testosterone | AST GGT Vitamin D BUN | Vitamin B12 ALT TSH Triglyceride Bilirubin total Iron Folate Lactate |
| SINGLE MEASUREMENT +/- range* | ~1-3% | ~5-7% | ~8-14% | ~15-25% | ~26-30% | ~40-50% |
| SERIAL MEASUREMENTS Change required** | ~2-5% | ~6-10% | ~11-20% | ~21-30% | ~35-45% | ~50-75% |

* based on the analytic and biologic variation

** also known as the reference change value

REVISED

Data collated primarily from here - <https://www.westgard.com/biodatabase1.htm>
but some taken and confirmed from a few other sources - numbers rounded off for ease of use
James McCormack BSc (Pharm), Pharm D - therapeuticseducation.org

If I was the boss of “LAB” result reporting

All of this could be done today

Shift from a laboratory perspective to a patient-centered viewpoint

Using **BALLPARK** estimates

ALWAYS provide the imprecision

Provide MUCH more definitive guidance

Stop using terms like low, medium, high or significant

If they are “risk factor” measurements then they should only be provided with “risk” estimates

Do not do a test without discussion of a pre-test probability and then provide a post-test probability

Make many tests more “inconvenient”?



As much as
humanely possible

DO NOT use “flags”, adjectives, or mention SDs, Gaussian distributions, two-sided tests, Z-scores, T-scores, confidence intervals, or p-values.

Not sure if we even need point estimates

Lab Value thoughts

have you first looked at how the patient is clinically doing?

will the result of your test change what you would do?

does a “risk factor” test improve your assessment of risk?

how big a change do you expect from your treatment?

what is the sensitivity and specificity of the test? - pre-test and post-test probability

how long does that change take?

how big a change is needed to be confident a change has occurred?



**When someone
does something
wrong, don't forget
all the things they
did right.**