

**James McCormack**  
**BSc (Pharm), PharmD**  
**Professor**  
**Faculty of Pharmaceutical Sciences**  
**University of British Columbia**  
**Vancouver, BC, Canada**

**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	Cervical Cancer	Food and Waterborne Illness	Inherited Copper Toxicity	Neural Tube Defects	Small Cell Lung Cancer
Acidosis and Alkalosis	CF	Food Poisoning	Insulin Resistance	Neuropathy	Spina bifida
Acidosis/Alkalosis	CFIDS	Fungal Infections	Insulin Resistance Syndrome	NHL	Spinal dysraphism
aCL Syndrome	CFS	Gastroenteritis	Iron Overload Disease	Non-Hodgkin lymphoma	Spinal Meningitis
ACS	CHF	Gluten-Sensitive Enteropathy	Iron Storage Disease	Non-Small Cell Lung Cancer	SSc
Acute DIC	Chlamydia	Gonorrhea	Jaundice	Nontuberculous Mycobacteria	Stable angina
Acute Idiopathic Polyneuritis	Chronic Fatigue and	Gout	JIA	Nontuberculous Mycobacteria Infections	Staph
Acute Inflammatory Demyelinating 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	Keratconjunctivitis 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	Syndrome 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 Adosteronism	Types of Liver Disease
Atypical Mycobacteria	Discoid Lupus	HL	Meningococcal Meningitis	Primary Hyperaldosteronism	Ulcerative Colitis
Atypical Pneumonia	Disseminated Intravascular Coagulation	Hodgkin Disease	Menopause	Prostate Cancer	Unstable angina
Autoimmune Diseases	Disseminated Intravascular Coagulopathy	Hodgkin Lymphoma	Metabolic Syndrome	Protein in urine	Urinary Tract Infection
Autoimmune Thyroiditis	Disseminated Lupus Erythematosus	Hospital-Acquired Pneumonia	MG	Proteinuria	UTI
Avian Flu	DJD	HPFH	MI	RA	Vaginal Infection
Bacillus anthracis infection	Double Pneumonia	HPV	Morphea	Reactive Arthritis	Vaginitis and Vaginosis
Bacterial Arthritis	Down Syndrome	Hughes Syndrome	MOTT	Reaven Syndrome	Vaginitis/Vaginosis
Bacterial Vaginosis	Drug-induced Lupus	Huntington Disease	MPDs	Renal Disease, Kidney Failure	Variant angina
Benign Prostatic Hyperplasia	DS	Huntington's Chorea Disease	MPNs	Rheumatoid Arthritis	Vasculitis
Benign Prostatic Hypertrophy	Dysmetabolic Syndrome	Hypercoagulable Disorders or States	MRSA	Rheumatoid Spondylitis	VD
Biological Warfare	Ebola Hemorrhagic Fever	Hyperparathyroidism	MS	Sarcoidosis	Venereal Diseases
Bioterrorism Agents	Ebola Virus Disease	Hypersensitivity	Multiple Myeloma	SCD	Vitamin B12 and Folate Deficiencies
Bleeding Disorders	Ebola Virus Infection	Hypertension	Multiple Sclerosis	Scleroderma	Vitamin B12 Deficiency
Blood in the urine	Encephalitis	Hyperthyroidism	Myalgic Encephalomyelitis	SEID	Vitamin K Deficiency
Bone Marrow Disorders	End Stage Renal Disease	Hypoparathyroidism	Myasthenia Gravis	Seizure Disorder	Vulvovaginitis
Borrelia burgdorferi Infection	Endocrine Syndromes	Hypothyroidism	Mycobacteria other than tuberculosis	Septic Arthritis	Walking Pneumonia
Borrelia mayonii Infection	Endocrine System and Syndromes	IBD	Mycoses	Sexually Transmitted Diseases	West Nile Virus
BPH	Epilepsy	ICterus	Myelocle	Sexually Transmitted Infections	Wilson Disease
Breast Cancer	ESRD	Infectious Arthritis	Myelodysplasia	Shingles	WNV
CAH	EVD	Infectious Polyneuritis	Myelodysplastic Syndrome	Sicca Syndrome	Wound and Skin Infections
Cancer	Excessive Clotting Disorders	Infertility	Myelomeningocele	Sickle Cell Anemia	
Candidiasis	Extracosseous Plasmacytoma	Inflammatory Bowel Disease	Myeloproliferative Disorders	Sickle Cell Disease	
Carbohydrate Intolerance	Fibromyalgia	Influenza	Myeloproliferative Neoplasms	Sjögren Syndrome	
Cardiovascular Disease	Flu	Influenza A	Mycardial Infarct	SLE	
Celiac Disease	Folate Deficiency	Influenza B	Neonatal Lupus		
Celiac Sprue	Folic Acid or B9 Deficiency	Inhalation anthrax	Nephrotic 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





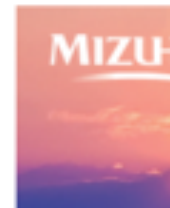
If you're not sick you just  
haven't had enough tests  
*Robert Rangno, MSc, MD*

# 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, %
<b>Preanalytical</b>		
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
<b>Analytical</b>		
Instrument-caused random error	3	1.9
Analytical inaccuracy not recognized	21	13.1
Subtotal	24	15
<b>Postanalytical</b>		
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

CVA - analytical variation

Biological Variation

CVI - within subject

CVG - between subject

Reference change values (RCV)

Analytic  
variation

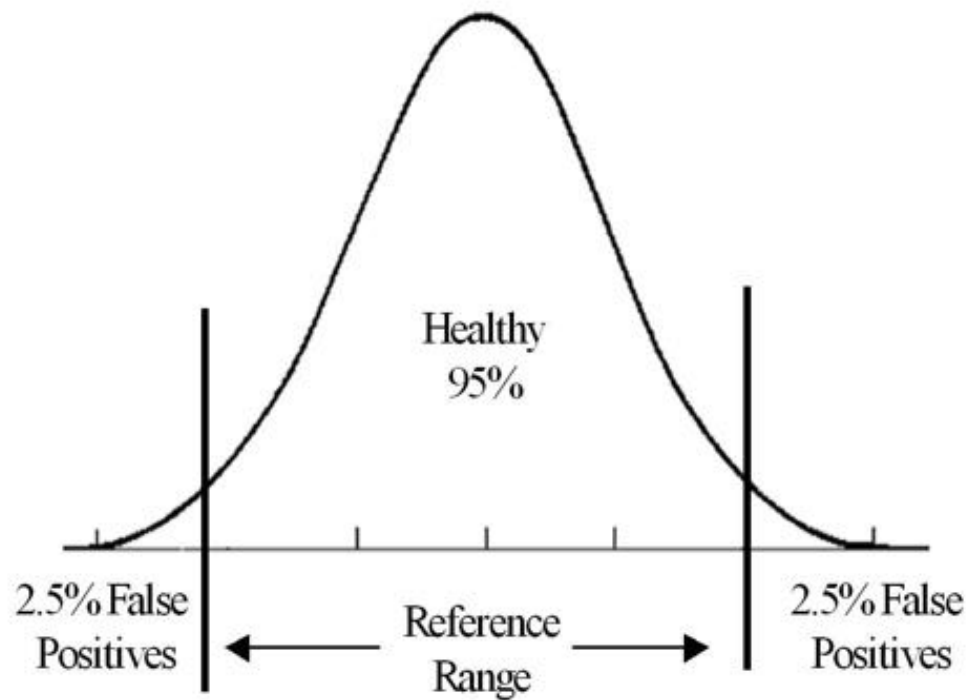
Biologic  
variation



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*

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

# Nerd Alert



# Reference Change Values (RCV)

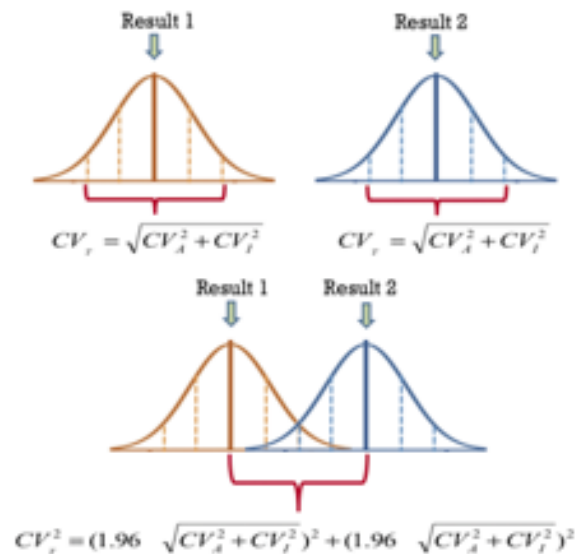
a tool for assessment of the significance of differences  
in serial results from an individual



# 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/10.1080/00036812.2016.1154108>



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

2014) and a statement on risk-limiting post-election audits

“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

**DEBATE**

**Open Access**

# How confidence intervals become confusion intervals

James McCormack<sup>1</sup>, Ben Vandermeer<sup>2</sup> and G Michael Allan<sup>3\*</sup>

## The Word SIGNIFICANT

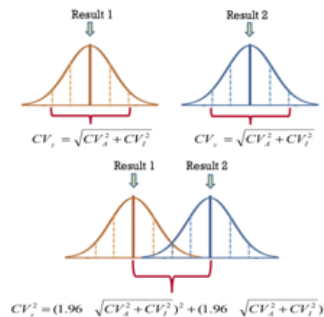


# Reference Change Values

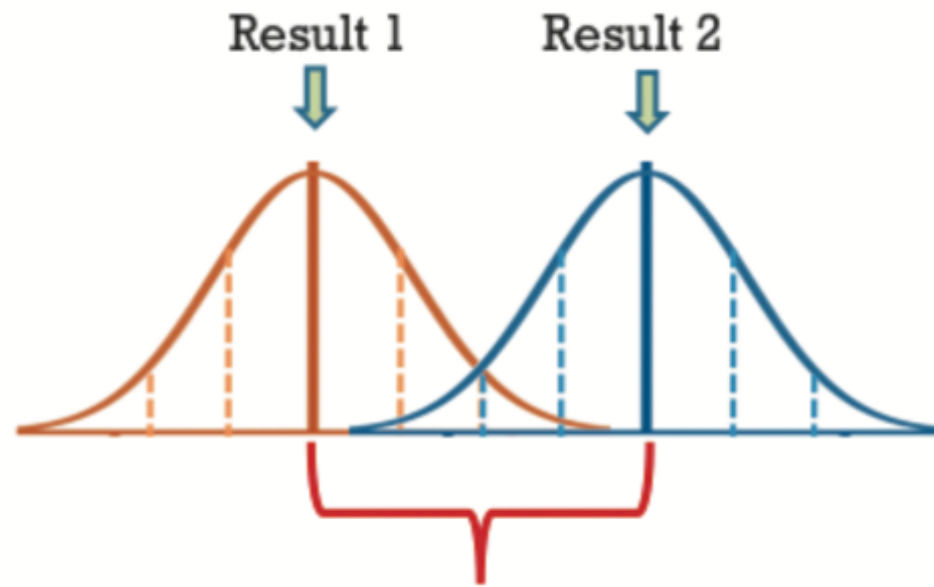
findings of a “significant difference” JUST means we are ruling out that the difference seen is due to chance

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~~

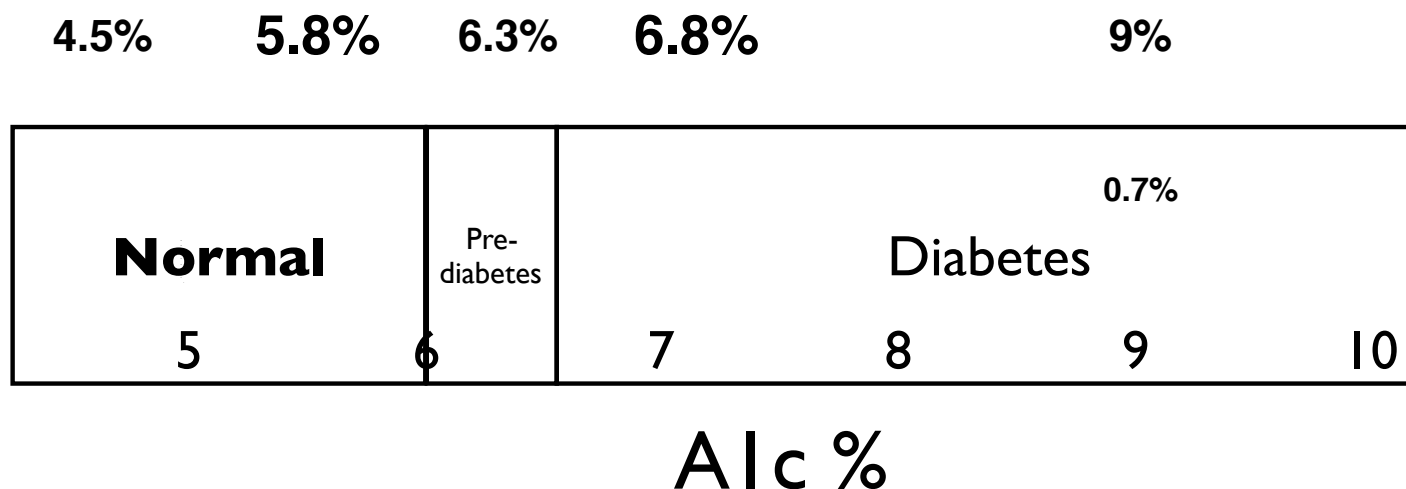




Glucose

# Precisely Imprecise

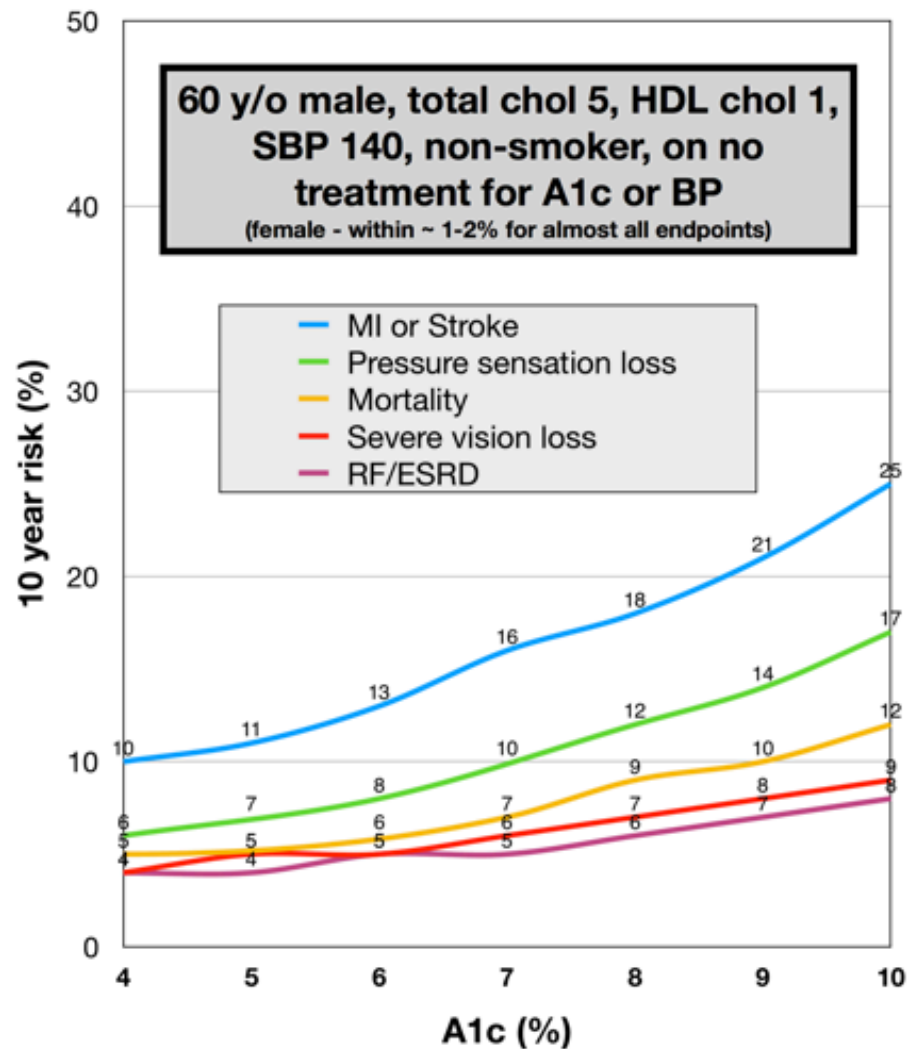
## What an A1c result really means



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



T2DM risk  
should not  
be  
categorized  
as  
YES  
or  
NO

<https://sanjaybasu.shinyapps.io/recodesi/> - from the ACCORD study





Cholesterol

## CHOLESTEROL

There are NO large studies that have looked at getting patients to different cholesterol levels

### **Effect of Lower Targets for Blood Pressure and LDL Cholesterol on Atherosclerosis in Diabetes**

The SANDS Randomized Trial

3 years - 499 American Indian men and women aged 40 years or older with type 2 diabetes and no prior CVD events

Results - surrogates improved - no change in clinical outcomes

JAMA 2008;299:1678-89

## **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 ( $\geq 19$  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

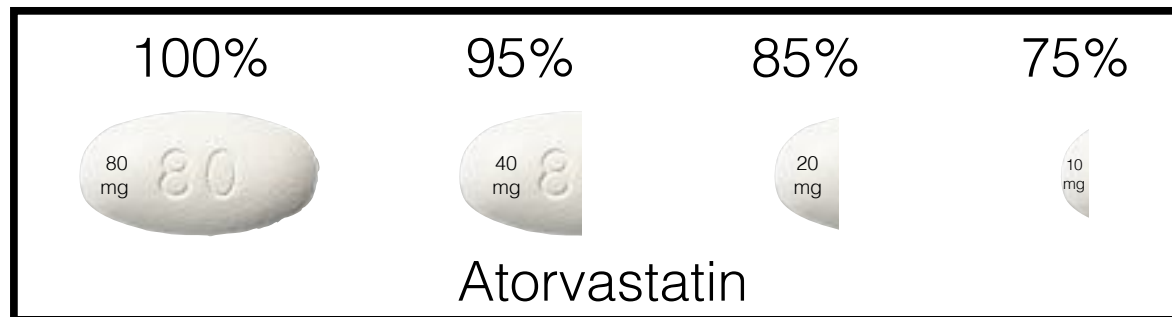
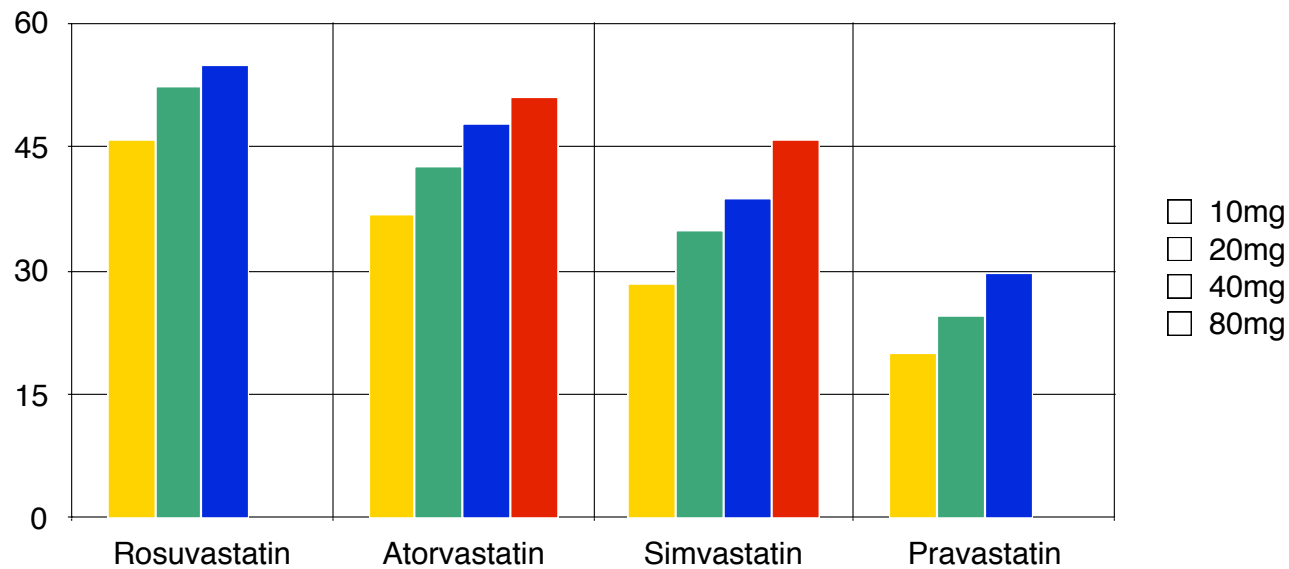
Total chol ~ - 0.80 to 0.80 mmol/L (~30 mg/dL)

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

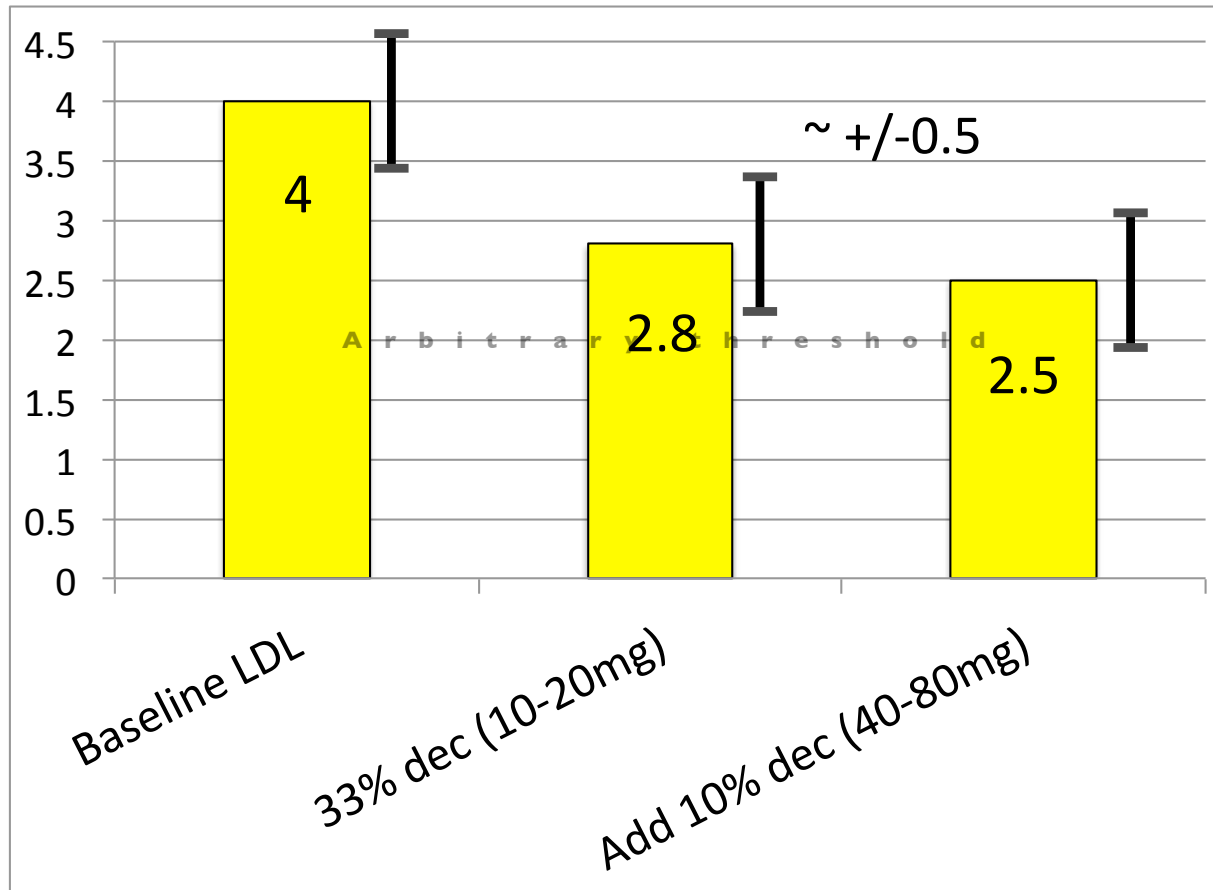
Average increase in cholesterol is 0.5-1%/year

“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



**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/TIA + peripheral vascular disease

---

**Age**

years

**Gender** ☐ Male ☒ Female

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

**Diabetes** ☐ Yes ☒ No

**Systolic Blood Pressure**

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**

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**

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.

[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
<b>NNT</b> ∞	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.](#)

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

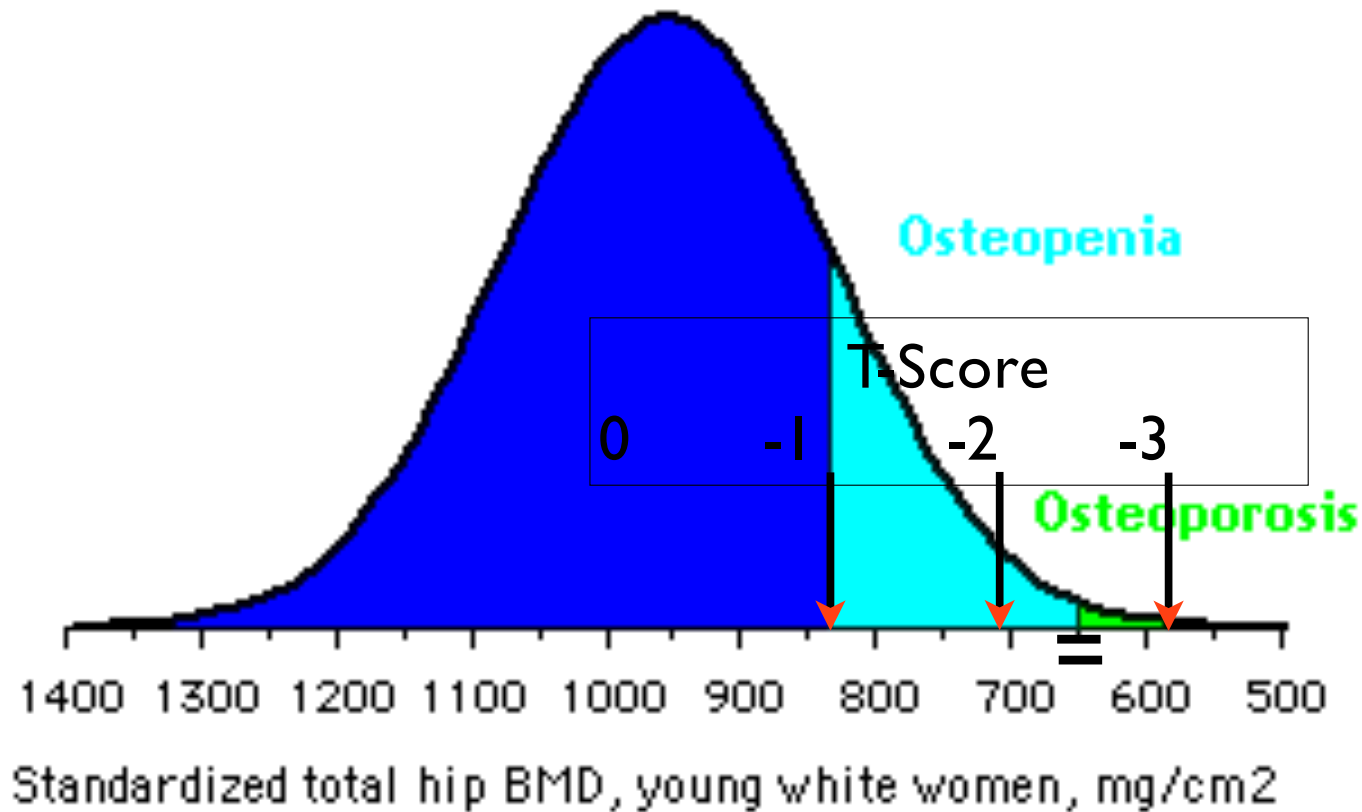
[cvdcalculator.com](http://cvdcalculator.com)





Bone density  
(almost all analytic issues)

- 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  $\pm 2-3\%$



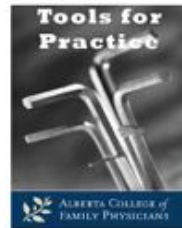
# Other Smarter People

## **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.”

Now What?!!



The Problem  
is  
NOT Fixable,  
it is  
Only  
KNOWABLE

TABLE - Approximate % Variance Estimates for Routinely Ordered Medical Measurements

	Analytical plus Biological	Analytical Variation	Analytical/Biological Variation
Test	+/- change required for 2 serial measurements (Reference Change Value)	+/- single measurement (Analytical CI)	+/- single measurement (Analytical + Biological CI)
Chloride	2-5%	1-2%	2-5%
Sodium	2-5%	1-2%	1-2%
Osmolality	2-5%	1-2%	2-5%
Bone Density (spine, total hip)	2-5%	1-2%	1-2%
Hemoglobin	6-10%	1-2%	6-10%
Bone Density (femoral neck)	6-10%	2-5%	2-5%
Calcium	6-10%	2-5%	6-10%
Protein	6-10%	2-5%	6-10%
PTT	6-10%	2-5%	6-10%
Albumin	6-10%	2-5%	6-10%
Potassium	11-20%	1-2%	6-10%
Magnesium	11-20%	2-5%	6-10%
PCO2	11-20%	2-5%	6-10%
A1c	11-20%	6-10%	6-10%
Glucose	11-20%	2-5%	11-20%
Creatinine	11-20%	2-5%	11-20%
ALP	11-20%	6-10%	11-20%
INR	21-30%	6-10%	11-20%
LDL cholesterol	21-30%	2-5%	11-20%
HDL cholesterol	21-30%	1-2%	11-20%
Total cholesterol	21-30%	1-2%	11-20%
LDH	21-30%	1-2%	11-20%
Uric acid	21-30%	1-2%	11-20%
Phosphate	21-30%	2-5%	11-20%
Rheumatoid factor	21-30%	2-5%	11-20%
Testosterone	21-30%	6-10%	21-30%
GGT	35-40%	2-5%	21-30%
Urea	35-40%	2-5%	21-30%
AST	35-40%	6-10%	21-30%
VR D	35-40%	6-10%	21-30%
VR B12	>50%	21-30%	35-40%
ALT	>50%	6-10%	35-40%
TSH	>50%	6-10%	35-40%
Triglyceride	>50%	2-5%	35-40%
Total bilirubin	>50%	2-5%	40-45%
Iron	>50%	2-5%	>50%
Lactate	>50%	2-5%	>50%
Folate	>50%	21-30%	>50%

# 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

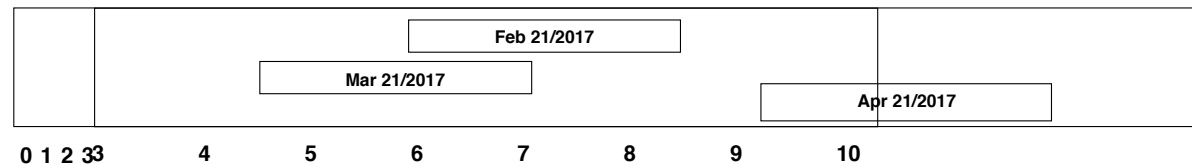
## Report Details

Close

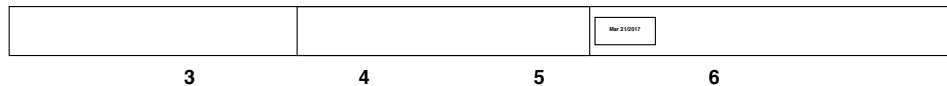
Patient Information				<input type="checkbox"/> Show More Patient Details	
Patient:	LLOYD, TESSA JANE ELSWITHA				
PHN:	9083126918	Phone:	(604)538-7132	Lab No:	17-141460231
		Flags Results	Reference Range	Units	
General Comments		Because of lab test and human variation, lab tests results can only be provided within a range			

**YOUR TRUE RESULTS ARE LIKELY SOMEWHERE IN THE RANGE INDICATED BY THE YELLOW STRIP**

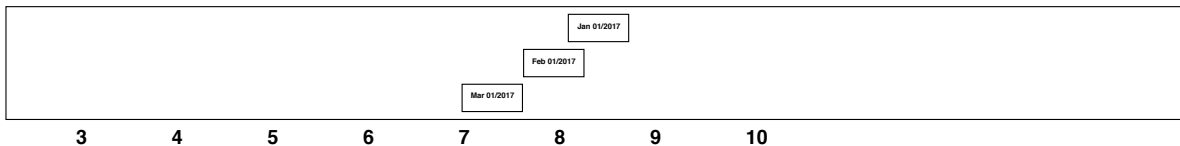
White blood cells



Potassium (mEq/L)



HbA1c (%)



95% of HEALTHY people have results somewhere in this range

5% of HEALTHY people have results somewhere in this range

Typically, results in this range are considered in the "normal" range

Typically, results in this range are considered outside of the "normal" range

There is no "normal/abnormal" range - the higher the value the higher your CVD risk - discuss your risk with your HCP

DATE 1

Typically, if 2 test results from different times overlap we consider the results to not be different

DATE 2

If they don't overlap then we likely think they have changed

<input type="text"/>	Deals with the reference interval	
<input type="text"/>	Deals with the analytical and human variability	DATE 1
<input type="text"/>	Deals with risk lab tests	DATE 2

OVERLAP Deals with reference change values

# 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?



Questions?

A black marker is shown drawing a curved line underneath the word 'Questions?' on a piece of lined paper.

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