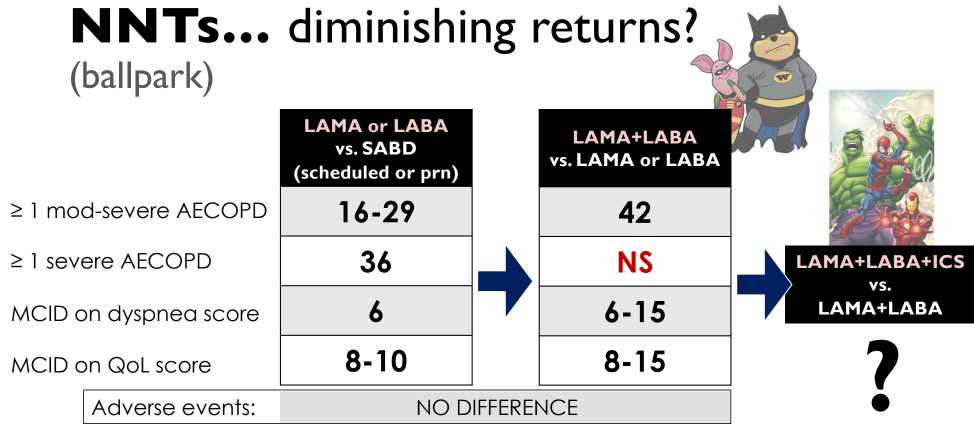


Figure 2. COPD Pharmacotherapy.



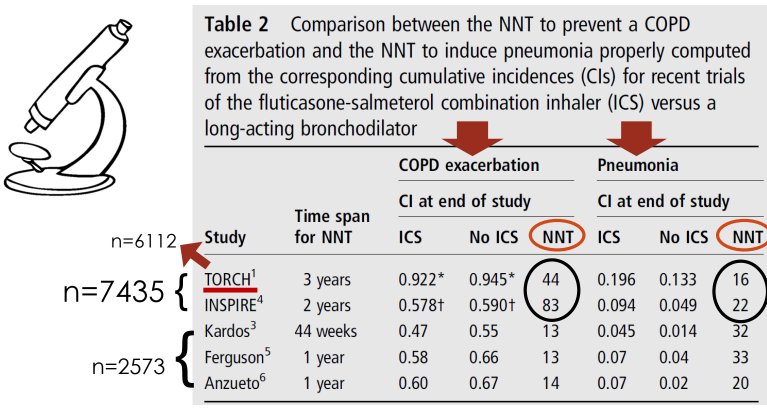
(how well is this collected and reported?)

Thorax 2016;71:15–25 Int J COPD 2017:12 907–922 CDSR 2018, Issue 12. Art. No.: CD012620

Respir Res 2017;18:196

COPD: What to Do with all These New Inhalers? Dalhousie CPD Academic Detailing Service, 2017

BACK IN TIME... AECOPD VS. PNEUMONIA IN CONTEXT



Suissa S. Thorax 2013;68:540-543.

LABA+ICS GOES DOWN IN FLAMES?

ORIGINAL ARTICLE

Indacaterol–Glycopyrronium versus Salmeterol–Fluticasone for COPD



N Engl J Med 2016;374:2222-34

- Patients (n=3362)
 - At least one moderate exacerbation in the past year
 - 75% were GOLD stage D (i.e. severe)

Results:

- 0.21 less exacerbations/pt/yr for LAMA+LABA
- 1.8 point difference in QoL (SGRQ)
- ¼ puff less/day of rescue inhaler
- Pneumonia: NNH = 63 for LABA+ICS

So, LAMA+LABA modestly better than LABA+ICS in the highest risk patient, and safer

NOTABLE TRIALS:

OPTIMAL
WISDOM
SUNSET n=6,630
TRIBUTE
KRONOS
IMPACT n= 6,221

N Engl J Med 2018;378:1671-80

Once-Daily Single-Inhaler Triple versus Dual Therapy in Patients with COPD

WHO? FEV1 = 45%, ≥ 1 AECOPD/yr (55% had ≥ 2)

WHAT? LABA+LAMA+ICS (fluticasone) vs. LABA+LAMA vs. ICS+LABA

What did they find @ 1yr?

- → mod-severe AECOPD = 0.3/pt/yr
- → I hospitalizations = 0.06/pt/yr
- → **I** mortality = **0.83%**, **NNT=120**

Did patients **FEEL BETTER**? → well...

- \rightarrow SQRQ -1.8 \rightarrow NNT MCID = 13
- → TDI change not reported → MCID NNT = 17?

IMPACT? \rightarrow yes, a bit \rightarrow

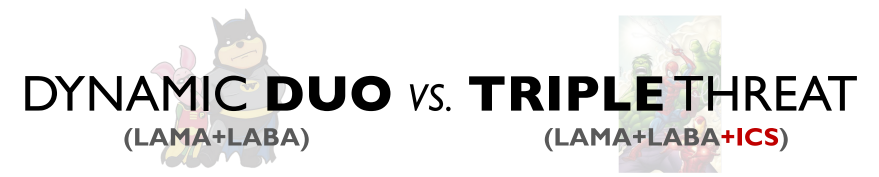
What's the CATCH?

you could have history of ASTHMA

• >70% on ICS pre-randomization

NNH (pneumonia) = 34





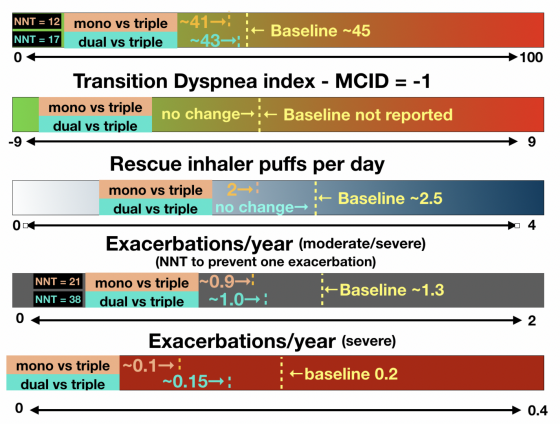
3 meta-analyses:

- Reduction in AECOPD (Cazzola, Eur Resp J 2018)
 NNT = 39 (for triple)
- Increase in PNEUMONIA (Zheng, BMJ 2018; Zayed, Clin Respir J 2019)
 NNH = 38-39 (against triple)

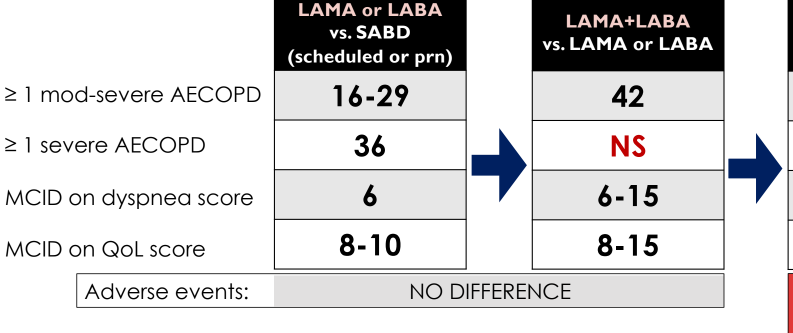
But, did they at least feel better day-to-day?

Ballpark estimates of the benefits seen from inhalers on clinically important outcomes

St George's Respiratory Questionnaire - MCID = - 4
(NNT to reach MCID)







any real net benefit?

LAMA+LABA+ICS vs. LAMA+LABA

38

0.05 less/pt/yr

NA

17

Pneumonia:

39

Thorax 2016;71:15-25 Int J COPD 2017:12 907-922 CDSR 2018, Issue 12. Art. No.: CD012620

Respir Res 2017;18:196

COPD: What to Do with all These New Inhalers? Dalhousie CPD Academic Detailing Service, 2017

ETHOS

N Engl J Med June 24, 2020;383:35-48

Triple Inhaled Therapy at Two Glucocorticoid Doses in Moderate-to-Very-Severe COPD

WHO? FEV1 = 43%, ≥1 AECOPD/yr (57% had ≥2)WHAT? LABA+LAMA+ICS (budesonide 320mcg or 160mcg) vs. LABA+LAMA vs. ICS+LABA

What did they find @ 1yr?

- \rightarrow \blacksquare mod-severe AECOPD = 0.35/pt/yr (or \sim 1 saved in 3 yrs)
- → **I** hospitalizations = **NS**
- $\star \to \blacksquare$ mortality = **1.0% NNT=100** (320mcg), 0.47%

Did (SUNE # 2) PEEE LOBETTER? → well...

- \rightarrow SQRQ change -1.9 (320mcg), -1.5 (160mcg) \rightarrow NNT MCID = 13-15
- → TDI change 0.4 (both doses) @24 wks → MCID NNT not reported

OF OTHER TRIPLE
TRIALS... VERY SIMILAR

What's the CATCH?

#1

you could have history of **ASTHMA**

80% on ICS pre-randomization 🜟

NNH (pneumonia) = 59



IMPACT: EFFECT OF ICS USE AT BASELINE ON AECOPD

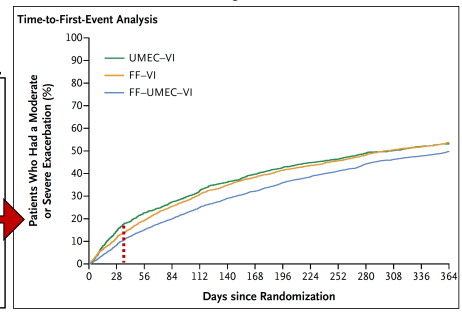
Am J Respir Crit Care Med;101(12):1508–1516, Jun 15, 2020 **Table 3.** Rates of On-Treatment Moderate/Severe Exacerbations in IMPACT by Medication at Study Entry

Baseline Medication*	FF/UMEC/VI (95% CI)	FF/VI (95% CI)	UMEC/VI (95% CI)
Overall ICS/LAMA/LABA ICS/LABA LAMA/LABA LAMA	0.91 (0.87–0.95) 1.21 (1.13–1.28) 0.70 (0.64–0.77) 0.84 (0.73–0.98) 0.65 (0.54–0.78)	1.07 (1.02–1.12) 1.43 (1.35–1.53) 0.85 (0.78–0.92) 1.11 (0.95–1.29) 0.75 (0.64–0.89)	1.21 (1.14–1.29) *\(\) 1.72 (1.58–1.87) 0.94 (0.83–1.06) 1.05 (0.86–1.29) 0.61 (0.47–0.80)

"...more than 70% were receiving an ICS, and patients with a history of asthma were included. Thus, for the patients assigned to the LAMA–LABA group, many of whom were actually stepping down in their treatment, ICS were abruptly withdrawn at the time of randomization... This design peculiarity, compounded by the probable inclusion of some patients who could have met a standard case definition of asthma, could explain the rapid surge in exacerbations observed in the first month after randomization in the LAMA–LABA group; during the subsequent 11 months of follow-up, the incidence of exacerbation with LAMA–LABA was practically identical to that with triple therapy."

Suissa, Drazen, NEJM April 18, 2018 NEJM

IMPACT trial: N Engl J Med 2018;378:1671-80



ETHOS & IMPACT: EFFECT OF ICS USE AT BASELINE ON MORTALITY

ETHOS

BGF 320/18/9.6 µg vs GFF Patients with an event, n (%) **BGF** 320/18/9.6 µg 18/9.6 µg Subgroup (N=2137) (N=2120) HR (95% CI) P-value 18/942 (1.9) 22/909 (2.4) 0.3539 1 moderate/severe exacerbation 0.74 (0.39, 1.40) ≥2 moderate/severe exacerbations 12/1195 (1.0) 34/1211 (2.8) 0.36 (0.19, 0.70) 0 severe exacerbations 23/1687 (1.4) 43/1691 (2.5) 0.51 (0.31, 0.85) ≥1 severe exacerbation 7/450 (1.6) 13/429 (3.0) 0.52 (0.21, 1.30) 0.1634 Post-bronchodilator FEV, <50 % predicted 27/1522 (1.8) 44/1522 (2.9) 0.62 (0.38, 0.99) 0.0468 Post-bronchodilator FEV, ≥50 % predicted 2/613 (0.3) 12/596 (2.0) 0.16 (0.04, 0.72) 0.0171 Triple therapy at screening 11/983 (1.1) 32/979 (3.3) 0.31 (0.15, 0.63) 0.0013 No triple therapy at screening 19/1154 (1.6) 24/1141 (2.1) 0.78 (0.43, 1.42) 22/1696 (1.3) 51/1698 (3.0) ICS at screening 0.41 (0.25, 0.69) 0.0006 No ICS at screening 8/441 (1.8) 5/422 (1.2) 1.49 (0.49, 4.55) 0.25 0.5 1.0 2.0 4.0 Favors triple therapy Hazard ratio

IMPACT

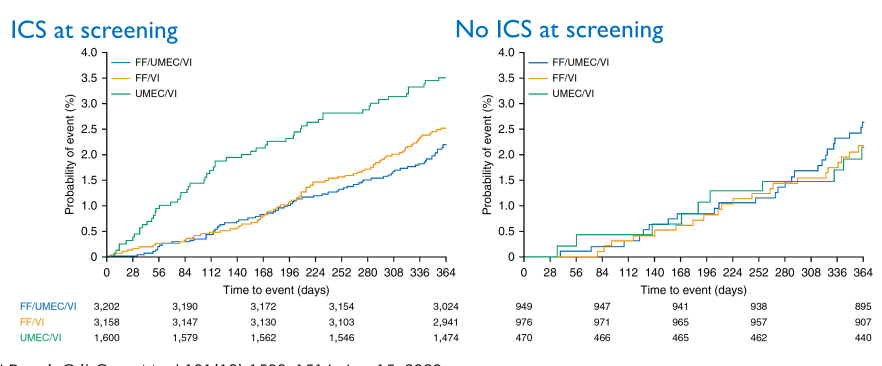
			Favors FF/UMEC/VI		Favors UMEC/VI		
	Patients with an event, n/N (%)					→	
Overall	FF/UMEC/VI	UMEC/VI				Hazard ratio (95% CI)	p-value
On-treatment ACM	50/4,151 (1.20)	39/2,070 (1.88)	-			0.58 (0.38, 0.88)	0.011
ACM including off-treatment data (with additional vital status follow-up) ICS therapy at screening	98/4,151 (2.36)	66/2,070 (3.19)	H			0.72 (0.53, 0.99)	0.042
On-treatment ACM	33/3,202 (1.03)	34/1,600 (2.13)	₩			0.44 (0.27, 0.71)	< 0.001
ACM including off-treatment data (with additional vital status follow-u p) No ICS therapy at screening	72/3,202 (2.25)	56/1,600 (3.50)	→			0.63 (0.44, 0.89)	0.009
On-treatment ACM	17/949 (1.79)	5/470 (1.06)	-	•		1.49 (0.55, 4.06)	0.430
ACM including off-treatment data (with additional vital status follow-up)	26/949 (2.74)	10/470 (2.13)	+	•	^	1.25 (0.60, 2.59)	0.550
Triple therapy at screening On-treatment ACM	13/1,672 (0.78)	15/864 (1.74)	→			0.40 (0.19, 0.84)	0.016
ACM including off-treatment data (with additional vital status follow-up)	36/1,672 (2.15)	30/864 (3.47)	-			0.62 (0.38, 1.00)	0.051
No triple therapy at screening							
On-treatment ACM	37/2,479 (1.49)	24/1,206 (1.99)	- →-			0.67 (0.40, 1.12)	0.129
ACM including off-treatment data (with additional vital status follow-up)	62/2,479 (2.50)	36/1,206 (2.99)	0 0.5 1.0	1.5 2.0	2.5 3.0 3.5	0.80 (0.53, 1.21) 4.0 4.5	0.285
				Hazard rat	io (95% CI)		

AJRCCM Articles in Press. Published November 30, 2020 as 10.1164/rccm.202006-2618OC

Am J Respir Crit Care Med;101(12):1508–1516, Jun 15, 2020



IMPACT: EFFECT OF ICS USE AT BASELINE ON MORTALITY



Am J Respir Crit Care Med;101(12):1508-1516, Jun 15, 2020

PRIMARY CARE vs. TRIALS

Plos One 2014;9(3):e90145

Table 2. Baseline comparison of the UNLOCK studies versus large COPD studies, including independent sample t-tests.

	-			
Characteristic	(primary care) UNLOCK studies	Large COPD studies (LPCS)	Mean difference between UNLOCK – LPCS (95% CI)	p-value
Patients (N)	3508	23860		
Age, years	66.1 (2.3)	63.7 (0.9)	-2.4 (-4.6 — -0.3)	0.03*
Male, %	60.9 (16.7)	73.3 (4.1)	12.4 (-3.1—27.9)	0.1
Current smokers, %	42.9 (9.5)	40.7 (8.6)	-2.2 (-13.2—8.8)	0.67
Pack years	43.6 (13.5)	44.9 (4.03)	1.3 (–15.2—17.8)	0.84
BMI, kg/m²	26.3 (0.5)	25.6 (0.9)	-0.7 (-20.6)	0.23
Postbronchodilator FEV ₁ , % predicted	63.8 (8.7)	47.4 (2.4)	-16.4 (-248.2)	<0.01*
FEV1:FVC, %	55.7 (0.7)	46.5 (4.0)	-9.2 (-14.1 —-4.2)	<0.01*
GOLD distribution				
Mild GOLD I	20.7 (13.2)	-	-	-
Moderate GOLD II	53.3 (6.2)	45 (6.3)	-8.3 (-16.6—0.1)	0.05
Severe GOLD III	21 (10.1)	44.5 (3.1)	23.5 (13.9—33.1)	<0.01*
Very severe GOLD IV	5.8 (5.2)	11.5 (3.5)	5.7 (-0.71—12)	0.08
Patient-reported outcomes				
SGRQ	32.6 (6.2)	48.4 (1.9)	15.8 (6.3—25.4)	0.01*
CCQ (mean)	1.6 (0.3)	-	-	-
MRC (mean)	2.1 (0.8)	2.7 (1.1)	0.6 (-1.5—2.7)	0.5
MRC score > 2 (%)	32.3 (17)	51.5 (2.1)	19.2 (1.3—37)	0.04*



https://www.trelegy.com

Better FEVI

Less GOLDBetter QoL

^{*} proportion of primary care patients eligible for inclusion in large RCTs \rightarrow 17% - 42%

THERE ARE A LOT OF "IFs": YOU GOTTA HAVE FAITH (OR HOPE)?

2 possible approaches:

1) PREVENTATIVE

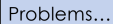
→ prescribe knowing that AECOPD are reduced overall

Keeping in mind...

- AECOPD occur relatively infrequently
- seasonal fluctuations not uncommon

2) SYMPTOM-based

→ prescribe the inhaler → assess if patient feels better



- Problems... COPD symptoms often fluctuate widely day-to-day/wk-to-wk (often > than differences in RCTs)
 - When are new inhalers started? → when patient feeling worse



