09/21/2020



UW Psychiatry & Behavioral Sciences

## Working in Primary Care Settings: How can I communicate effectively with PCPs?

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## **Speaker Disclosures**

• Dr. Kern has no disclosures to make.

Integrated Care Training Program

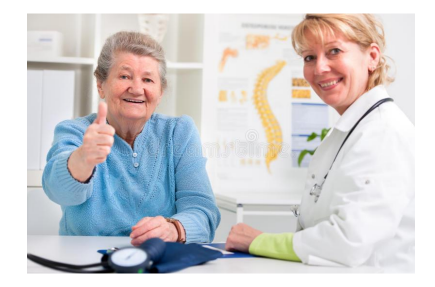
## **Learning Objectives**

At the conclusion of this presentation, the learner will be able to:

- 1. Describe the central role of the primary care provider in the collaborative care team.
- 2. List three strategies to improve the effectiveness of communication with your primary care provider partner.

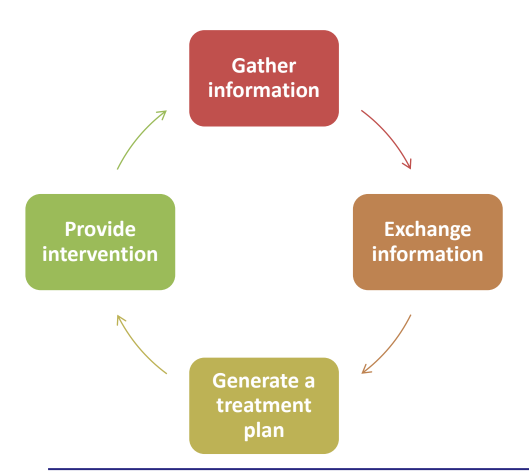
## Why the PCP is important

- PCP recommendation is powerful
  - Introduce care manager and team roles
- Existing relationship is foundation for alliance with the Collaborative Care team



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## **PCP Role: Diagnosis**

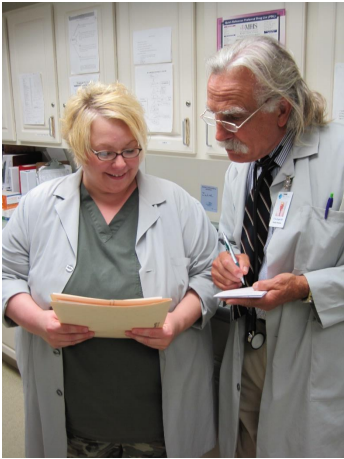


 PCP may have long history with patient

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# Engaging the PCP: "Why am I doing this?"

- These patients are already your patients.
- They are not going away.
- We can help with everyday workflow, shorten long appointments, reduce arguments about controlled substances... We have your back!
- Can help with chronic disease outcomes, IMPROVE YOUR METRICS!



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## Making yourself indispensable



- Respond to "extra" requests
- Make sure you are "interruptible"
- Point out that you can respond to patients that take large amounts of PCP time. Help them develop the skill of quick and effective referral.

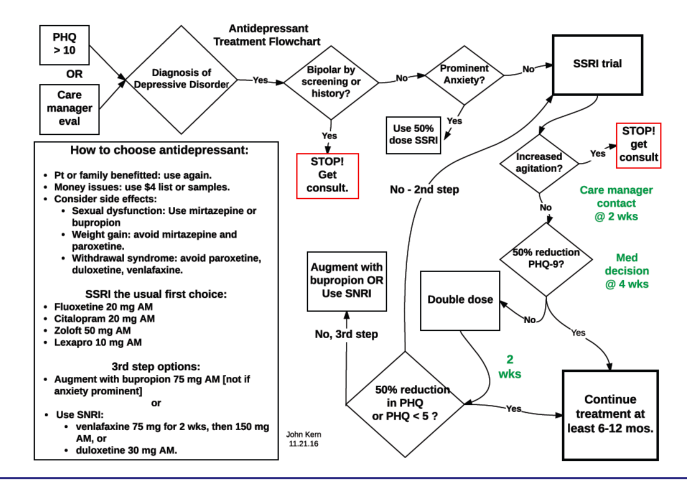


# Making yourself indispensable some examples:

- Treatment protocols an evolving toolkit
  - Medication info
  - Depression
  - ADHD
  - Bipolar
  - Sleep
  - Smoking, other behavior change
- Practical help with managing difficult patients
  - Benzos
  - Pain
  - Suicide risk
  - MAT?

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## Example of how to come bearing gifts: antidepressant protocol



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## **Prescribing Cheat Sheet**

NAME Generic (Trade)	DOSAGE	Key Clinical Information
	Antide	pressant Medications*
Bupropion (Wellbutrin)	Start: IR-100 mg bid X 4d then $\uparrow$ to 100 mg tid; SR-150 mg qam X 4d then $\uparrow$ to 150 mg	Contraindicated in seizure disorder because it decreases seizure threshold; stimulating; not good for treating anxiety disorders; second Ine TX for APDA source potential, 6 (IRSR), S. OL)
Citalopram (Celexa)	bid; XL-150 mg qam X 4d, then 1 to 300 mg qam. Range: 300-450 mg/d. Start: 10-20 mg qday, 110-20 mg q4-7d to 30-40 mg qday. Range: 20-60 mg/d.	ume i A tor AUFU, acouse protental. € (HOSH), § (AL) Best tolerated of SSRis; very few and limited CVP 450 interactions; good choice for anxious pt. €
Duloxetine (Cymbalta)	Start: 30 mg gday X 1 wk, then 1 to 60 mg gday, Range: 60-120 mg/d.	best oberated at Sarrays, very few and namedo the resonance accurs, good choice for anticolas pr. § More Gl side effects than SSRs: to neuropartitic pair, need to monitor BPS, 2 <sup>od</sup> in the for ADHD, §
Escitalopram (Lexapro)	Start 5 mg qday X 4/7d then 1 to 10 mg qday. Range 10-30 mg/d (3X potent vs. Celexa).	Best tolerated of SSRIs, very few and limited CYP 450 interactions. Good choice for anxious pt. \$
Fluoxetine (Prozac)	Start: 10 mg gam X 4-7d then ↑ to 20 mg gday. Range: 20-60 mg/d.	More activating than other SSRIs; long half-life reduces withdrawal (t ½ = 4-6 d). ¢
Mirtazapine (Remeron)	Start 15 mg qhs. X 4-7d then 1 to 30 mg qhs. Range: 30-60 mg/qhs.	Sedating and appetite promoting; Neutropenia risk (1 in 1000) so avoid in immunosupressed patients. ¢
Paroxetine (Paxil)	Start 10 mg qhs X 4-7d then 1 to 20 mg qday Pange: 20-60 mg/d.	Anticholinergic; sedating; significant withdrawal syndrome. ¢
Sertraline (Zoloft)	Start: 25 mg qam X 4-7d then ↑ to 50 mg qday. Rn 50-200 mg/d.	Few and limited CVP 450 interactions, mildly activating, é
Venlafaxine (Effexor)	Start IR-37.5 mg bid X 4d then ↑ to 75 mg bid; XR-75 mg 4then ↑ to 150 qAM. Range: 150-375 mg/d.	More agitation & Gl side effects than SSRIs; tx neuropathic pain above 150 mg qday; need to monitor BP; 2 <sup>rd</sup> line tx for ADHD. Significant withdrawal syndrome. ¢ (IR). S.OCR).
"Warnings/precautions: 1) Po SSRIs and SNRIs), Increased	stential increased suicidality in first few months, 2) Long term weight gain d risk of bleeding with SSRIs and SNRIs (especially in combo with NSAIDs)	roptont, 3) Sexual side effects con tome (except bupmpion), especia
	A	Includes information such as:
Alprazolam (Xanax)	Start: 0.25 mg – 0.5 mg tid. Usual MAX: 4 mg/d.	
Chlordiazepoxide (Librium)	Start: 10-20 mg 3-4X daily. Usual MAX: 200 mg/d	
Clonazepam (Klonopin)	Start: 0.25 mg bid or tid. Usual MAX: 3 mg/d.	<ul> <li>Basic education</li> </ul>
Diazepam (Valium)	Start: 2–10 mg bid to gid with doses depending on symptoms severity. Usual MAX: 30-40 mg/d.	
Lorazepam (Ativan)	Start: 0.5-1 mg bid to tid. Usual MAX: 6 mg/d. Insomnia: 0.5-2 mg qhs.	Equiv. dose?
Buspirone (Buspar)	Start 7.5 mg bid. Range: 10-30 mg bid.	Non-berzo SSR-Illa. Arthistamic-antenentb.
Hydroxyzine (Vistaril)	Start: 25-100 mg 3-4 X per day. Usual MAX: 400 mg per day. Start: 1 mg ghs. Increase g 2-3 d until symptoms abate. Usual MAX: 10 mg ghs.	Of antipyetensive used a Names and doses of
Prazosin (Minipress)	otare ring quo, indease q 2 o d'ana symptonis asate, osata nexe, roing quo.	after each average active de la
Trazodone (Desyrel)	Start: 25-50 mg qhs. Range: 50-150 mg/qhs.	Commonly used as sleep air
Temazepam (Restoril) Zolpidem (Ambien)	Start 15 mg at bedtime. MAX: 45 mg qhs.	Ty: 2 & https://dot.initia.com/
Zoipidein (Ambien)	Start 5-10 mg qhs. MAX: 20 mg qhs.	
		Mood Stabilizers medication
1.144-1	Start: 300 mg bid to tid. Target plasma level: acute mania & bipolar depression: 0.8-1.2 mea/L: Maintenance: 0.6-0.8 mea/L. Available in ER form dosed once daily (usually at HS.	Black box warning for tool
Lithium	Lithobid & Eskalith). Plasma levels related to renal clearance.	I SH and SMP before statun Clearance. Lihimu strongly a
	Start 750 mg daily (bid or tid, DR; gday, ER); increase dose as quickly as tolerated to	
Divalproex (Depakote)	clinical effect. Target plasma level: 75 to 100 mcg/mL (DR) & 85-125 mcg/ml (ER).	this risk). Need to monitor 1 • ( OMMON CIDO OTTOCTO
	Start: 25 mg daily for weeks 1 & 2, then 50 mg daily for weeks 3 & 4, then 100 mg qday for	
Lamotrigine (Lamictal)	week 5, and finally 200 mg qday for week 6+ (usual target dose). Dosage will need to be	2000). No drug level monto side effects. 6
	adjusted for patients taking enzyme-inducing drugs or Depakote.	
		EPS: moderate (especially a final data in the especial data in the espec
	Mania. Start: 15 mg qday; Range: 15-30 mg/day. MDD adj bc. Start: 2-5 mg/day; adjust	
Aripiprazole (Abilify)	dose q 1+ weeks by 2-5 mg. Range: 5-10 mg/day. MAX: 15 mg qday. Schizophrenia. Start: 10-15 mg/day: ↑ at 2 week intervals: rec. dose: 10-15/day. MAX: 30 mg/day	indication for adjunctive trea
Olemanias (Zeneral)	Start. 5-10mg daily titrating to 15-30 mg daily once or divided bid.	EPS: Low; Metabolic side ef
Olanzapine (Zyprexa)		lipids regularly. \$
	Bipolar Dep: Start: 50 mg qhs; Initial target: 300 mg qhs; Range: 300-600 mg/d Mania.	EPS: Lowest (except for Clozaril); Metabolic side effects: moderate. Highly sedating. FDA indication for bipolar depression and adjunctive
Quetiapine (Seroquel)	Start: 50 mg bid; Initial target: 200 mg bid. Range: 400-800 mg/d. MDD adj b:. Start: 50 mg ghs; Initial target: 150 mg ghs; Range: 150-300 mg/day. Schizophrenia, Start: 25 mg	treatment of MDD. Potential increased suicidatily in first few months. Need to screen glucose and lipids regularly. Abuse potential. Available in an extended release form: Screen usIR S, GIR & SR, Avoid or use alternative in combination with methadone due to OTE profongation. S
	bid and increase by 50-100 mg/d (bid/tid). Initial target: 400 mg/d. Range: 400-800 mg/d	an exercised release with services risk a rest, revised as an entrance in some national and interface to give providents.
Risperidone (Risperdal)	Start 0.5 – 1mg qhs or bid titrating to 4-6 mg daily or bid. Available as long-acting injectable given q 2 weeks called Risperdal Consta.	EPS: highest; Metabolic side effects: moderate. Hyperprolactinemia and sexual side effects common. Need to screen glucose and lipids regularly, c
Ziprasidone (Geodon)	Start: 40 mg bid titrating quickly to 60–80 mg bid. Needs to be taken w/ food (doubles	EPS: moderately high (especially akathisia); Metabolic side effects: lowest. Need to screen glucose and lipids regularly. Lower dosage can be
-	absorption).	more agitating than higher doses. Contraindicated in combination with methadone due to OTc profongations. So mis nederly patients with dementia: 2) Increased inside of OTc protogiation and risk of sudden death (respective) in combination with other drugs
**Antipsychotic/mood stabil that are known to prolong t		ins in energy patients with dementia, 2) moreased risk of QTC protongation and risk of sudden death (especially in combination with other drugs

https://aims.uw.edu/sites/default/files/Psychotropics%20Medications\_2018.pdf

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## **Bipolar Roadmap**

#### Bipolar Management Roadmap plagnosis History, including prior treatment MDQ, then CIDI if positive

#### Care Manager Consultation -

Confirm diagnosis Is specialty care needed? Consult with psychiatrist before making diagnosis, or changing treatment. Give Information Packet; Diagnosis Medication Info Mood Charting Rhythm / self-management / sleep hygiene

#### Arrange aftercare

No more than 2 wks with new of changed meds No more than 3 months ever Call for no show Follow mood charts.

#### How to choose mood stabilizer:

- If antidepressant on board, discontinue.
- Lithium first line. Usually Depakote 2nd, Tegretol 3rd.
- If psychotic atypical

John Kern 11.22.16

- If depressed: Lamictal / Latuda / Seroquel
- Not unusual to need more than one mood stabilizer.

#### Lithium:

Start 300-600 mg hs, titrate to response weekly and to level ~0.7. Lab monitoring:

> Baseline TSH, BMP, Lithium level at one week with each change, then q 6 mos with BMP when stable. TSH yearly

#### Side effect mgmt:

Tremor [lower dose or add proranolol 20 mg prn. Gl upset (divide dose, take with food.) Loose stools, acne, wt gain, <u>polyria</u>. Serious but rare: renal insufficiency.

#### Valproate

Start 20 mg/kg/day = weight in Use x 10 rounded to 500 mg. HS dosing Laboratory monitoring: CBC, CMP baseline, at <u>one month</u> Levels at one month, with dosage change, lack of efficacy. Target level: 50-120 Titrate to effectiveness. Side effect management: Weight gain - dietary management Tremor - beta-blocker Gi distress - bs dose Risk of PCOS - avoid in young women, rash Serious but rare: Hepatotoxicity [minor increase in LFT's is not unusual], encephalopathy, Pancreatitis, bone marrow d/o

#### Carbamazepine:

200 mg BID x 2 wks, then increase by increments of 200 mg per day as tolerated. Laboratory monitoring: level at one month, 3 months, with dosage change, lack of efficacy, side effects, watch for induction Target levels 4-12, cbc, & cmp at one month Side effect management: Ataxia - reduce dose Hyponatremia monitor, discontinue below Na 125. Rash Serious but rare: Stevens-Johnson syndrome Bone marrow disorders

#### Lamictal

Titrate per instructions: 25 mg daily x 2 wks, then 50 mg daily x 2 wks, then 100 mg daily. If on Depakote, 25 mg every other day x 2 wks, then 50 mg. May not need more than 25-50 mg. If on Tegretol, 50 mg daily x 2 wks, then 100 mg daily Labs - not recommended Side effect management:

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In the break room at your new clinic, you are introduced for the first time to a PCP with whom you will be working. She says, "Nice to meet you. I have five minutes until my next patient. What's up with this Collaborative Care thing?"

- What are three things you could say to help build your working relationship?
  - Explaining your role
  - How Collaborative Care is different from treatment as usual in primary care
  - How can you be useful to them

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## How the psychiatrist leads the team:

- Training and shaping care manager practice over team.
- Advocating for the program with administration.
- Improving practice via attention to data, quality improvement.
- Framing the significance of the team's function they don't know that they work at the cutting edge, they are just going to work.
- Point out all the advantages to psychiatry in primary care
  - Urgent access
  - Lab monitoring
  - Systematic approach to care

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## Takeaways

- 1. PCP engagement crucial to a successful Collaborative Care program.
- 2. Understanding needs and constraints of PCP goes a long way to engagement.
- 3. Ongoing curiosity about how to be more helpful to your PCP partner will inspire your creativity.

## Resources

- <u>AIMS Center office hours</u>
- <u>UW PACC</u>
- Psychiatry Consultation Line
   (877) 927-7924
- Partnership Access Line (PAL)
   (866) 599-7257
- PAL for Moms
  - (877) 725-4666

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## **Questions and Discussion**

 Ask questions in the chat or unmute yourself



## Registration

 If you have not yet registered, please email <u>uwictp@uw.edu</u> and we will send you a link

