#### Hepatitis C Health Summary

Name:		Pertinent Medical History:						
DOB:		Previous hepatitis C treatment						
Phone #:		Specify: Cirrhosis <sup>1</sup>						
<b>Alternate Contact</b>	<b>:</b>			⊔ NO				
Medications <sup>2</sup> :		Child-Pugh Score: Other Liver Disease <sup>1</sup>		- No				
ivieuications .								
		Specify: Pulmonary Disorders <sup>1</sup>						
				⊔ NO				
		Specify: Cardiac Disease <sup>2</sup>	□ Yes					
		Specify: DVT or PE <sup>1</sup>						
		Specify: PPI/H2 blocker/Antacid use <sup>2</sup>	□ Yes					
			⊔ res	⊔ NO				
		Specify: Autoimmune Disorders <sup>2</sup>	□ Yes					
		Specify:						
		Cancer	□ Yes					
		Specify: Current infection <sup>1</sup>						
		Specify:						
		High Blood Pressure	□ Yes					
		High Cholesterol	□ Yes	□ No				
		Kidney Disease <sup>2</sup>	□ Yes	□ No				
		Anemia <sup>1, 2</sup>	□ Yes	□ No				
		Current TB Treatment <sup>2</sup>		□ No				
		Diabetes Specify Type 1 or 2		□ No				
Allergies:		HIV or AIDS <sup>1</sup>	□ Yes	□ No				
		Seizure Disorder <sup>2</sup>	□ Yes					
		Depression/Anxiety						
		Other Psychiatric Conditions						
Labs Prior to Trea	tment:	Specify:		<b>.</b>				
	: □ Pregnancy test	Screen & Review: AUDIT-C	PHO-	-9				
miniculately prior	☐ Uric Acid (ribavirin only)	Vaccine Status (give if needed):						
Within 1 month:	□ CBC with differential	Hepatitis A (If unknown, che	eck hep A	total IgG)				
Within I month.	☐ CMP (If GFR <30, do not start tx ¹)	Hepatitis B (If unknown, che	•					
	□ PT/INR	Other vaccines as appropr	iate:					
	□ HCV RNA	<ul><li>Flu (annually)</li></ul>						
Within 3 months:	☐ Genotype confirmation	□ PCV-13 (≥ age 65 or immunosuppressed)						
	☐ HBV DNA (if HBV cAb or sAg +)	□ PPSV-23 (≥ age 50 AN						
Within 6 months:		☐ Td (once every 10 years) <b>OR</b> Tdap (once)						
	□ TSH	□ Zoster (≥ age 60)		,				
	☐ A1C or Fasting Glucose	<ul><li>□ ECG (over age 65 or h/o cardiac disease)</li><li>Birth Control: Birth Control Methods:</li></ul>						
	□ Vitamin D 250H							
Within 1 year:	□ HIV screening	Females: LMP: Pregr		s 🗆 No				
, 22	□ NS5A RAV (genotype 3 only)	Males: Is your partner pregnan	t? □ Ye	es 🗆 No				
Once:	☐ IL-28b (if considering 8 weeks)	□ Counsel about pregnancy pr	evention	(see				
	( 2 2 2 2 3 2 2 2 2 2 2 2 2 2 2 2 2 2 2	Treatment Agreement)						
		□ Hepatitis C Treatment Agree	ment rev	viewed and				
		signed						

- 1- Further evaluation as indicated; consult Liver Disease Specialist prior to treatment.
- 2- Check drug interactions to treatment drugs. Further evaluation as indicated.



Liver Disease & Hepatitis Program
4315 Diplomacy Drive, Anchorage, AK 99508
Phone: 907-729-1560 Fax: 907-729-1570

http://www.anthc.org/hep

Follow us on Twitter:

Liver Program @ANTHCLiver

We are glad to hear you are interested in treatment for hepatitis C!

Here are some things to think about (and do) before you make your final decision about treatment:

<u>Why do treatment now?</u> New medicines have increased the chance of cure and have fewer side effects.

**Some people have worse liver disease than others.** If you have more severe liver disease (a lot of scarring in the liver or cirrhosis) you should consider getting treatment sooner.

#### What will happen during treatment?

There are 6 FDA approved treatment options for **genotype 1**:

- Option 1 is Harvoni<sup>®</sup> (ledipasvir/sofosbuvir), 1 tablet taken once a day for 8-24 weeks. The most common side effects are feeling tired and headache. In clinical studies, treatment response rates to Harvoni<sup>®</sup> were 94-100%.
- Option 2 is Epclusa® (sofosbuvir/velpatasvir), 1 tablet taken once a day for 12 weeks. The most common side effects are headache and feeling tired. In clinical studies, treatment response rates to Epclusa® were 94-98% for genotype 1.
- Option 3 is Viekira Pak™ (ombitasvir, paritaprevir, and ritonavir tablets; dasabuvir tablets) co-packaged as 3 tablets in the morning and 1 tablet in the evening with food for 12-24 weeks. The major side effects are nausea, itching, and insomnia. In clinical studies, response rates to Viekira Pak™ treatments were 86-100%.
- Option 4 is Zepatier™ (elbasvir/grazoprevir), 1 tablet taken once a day for 12-16 weeks. The most common side effects are feeling tired, nausea, and headache. In clinical studies, treatment response rates to Zepatier™ were 95-100%.
- Option 5 is Olysio® (simeprevir) plus Sovaldi® (sofosbuvir) 1 tablet of each taken once daily for 12 weeks. The most common side effects are feeling tired, headache, and nausea. In clinical studies, treatment response rates to Olysio® and Sovaldi® were 86-100%.
- Option 6 is Daklinza™ (daclatasvir) plus Sovaldi® (sofosbuvir) 1 tablet of each taken once daily for 12 weeks. The most common side effects are headache and feeling tired. In clinical studies, treatment response rates to Daklinza™ and Sovaldi® were 50-100%.

The FDA-approved **Genotype 2** treatment is Epclusa® (sofosbuvir/velpatasvir), 1 tablet taken once a day for 12 weeks. The major side effects are headache and feeling tired. In clinical studies, the treatment response rate to Epclusa® was 99% for genotype 2.

There are 2 FDA-approved treatment options for **genotype 3**:

- Option 1 is Epclusa® (sofosbuvir/velpatasvir), 1 tablet taken once a day for 12 weeks.
   The most common side effects are headache and feeling tired. In clinical studies, treatment response rates to Epclusa® were 85-98% for genotype 3.
- Option 2 is Daklinza™ (daclatasvir) and Sovaldi® (sofosbuvir) 1 tablet of each taken once daily for 12 weeks. The most common side effects are headache and feeling tired. In clinical studies, treatment response rates for Daklinza™ and Sovaldi® were 58-98%.

Some treatments will require ribavirin which is 5-6 additional tablets divided between morning and evening with food. The major side effects are feeling tired, nausea, itching and skin rash, trouble sleeping, irritability and weakness. A common side effect of ribavirin is anemia.

PLEASE NOTE: Ribavirin cannot be given to a pregnant or breastfeeding female or to a female who plans to become pregnant <u>or</u> a male who plans to father a child during or for 6 months after treatment because it can cause birth defects. There are no studies on Harvoni®, Epclusa®, Sovaldi®, Viekira Pak™, Zepatier™, or Daklinza™ in pregnant women or nursing mothers. Safety/risk during pregnancy or breastfeeding has not been established.

#### Are you ready for treatment?

To ensure that you will be successful in completing hepatitis C treatment we ask that the following items be done before starting treatment. We will review them together.

- You must be alcohol and drug-free. If you have recent drug/alcohol abuse, you need to be in an approved drug treatment program.
- You need to discuss hepatitis C treatment with your primary care provider and get his or her "OK" to start treatment.
- You should have a relative/close friend who is willing to help support you during treatment.
- You need to be committed to making every treatment appointment and getting **FREQUENT** blood draws (every 1-4 weeks). We will want to follow you very closely during treatment.

#### Additional Requirements If Checked:

months). This ultrasound checks your liver for cancer.

If yo	ou have cirrhosi	is, you may	y need an	EGD	(when	a docto	or loo	ks into	your	esop	hagu	JS
and stomac	h for swollen ve	eins that ca	ın bleed).									
If yo	ou have cirrhos	is, you nee	ed to have	e an	ultrasou	nd of	the liv	ver (do	ne in	the	past	6

Once everything you need to do on the list has been done, call your primary care provider to make an appointment to plan for hepatitis C treatment. At this appointment, treatment and side effects will be discussed in detail.

If you are coming to Anchorage and want a Fibroscan, call the Liver Clinic ahead of your visit to schedule. Fibroscan is a test using ultrasound waves to check liver stiffness or scarring/fibrosis in your liver. Fibroscan testing is done in the Internal Medicine Clinic. Do not eat or drink for 3 hours before the test.

### Congratulations on completing all the pre-treatment requirements!

**Hepatitis C Treatment Checklists** 

	ricpatitis e ricat	inche Checkhata					
Prior to Treatment							
Labs							
Immediately prior:	Pregnancy test (if applicable) Uric Acid (with ribavirin )	Miscellaneous Henatitis	: A (If vaccine status is				
Within 1 month:	CBC with differential	unknown, draw HAV total)					
Within I month.	CMP <sup>1</sup>	Hepatitis B (If vaccine status is					
	<del></del>		-				
	PT/INR		wn, draw HBsAg & HBsAb)				
	HCV RNA	PHQ-9 bas	seline				
Within 3 months:	Genotype confirmation	AUDIT-C					
	HBV DNA (if HBV cAb or sAg +)		bout pregnancy prevention				
Within 6 months:	AFP	Review &	sign Treatment Agreement				
	TSH						
	A1C or Fasting Glucose						
	Vitamin D 25OH (treat if deficient)						
Within 1 year:	HIV screening						
, , , , , , , , , , , , , , , , , , , ,	NS5A RAV (genotype 3 only)						
Once:	IL-28b (if considering 8 weeks)						
Office.	IL-28b (II considering 8 weeks)						
8 week	12 week	16 week	24 week				
Week 4	Week 2 (with ribavirin)	Week 2 (with ribavirin)	Week 2 (with ribavirin)				
HCV RNA	CBC	CBC	CBC				
CBC <sub>1</sub>	$\underline{\hspace{1cm}}$ CMP <sup>1</sup>	CMP <sup>1</sup>	CMP <sup>1</sup>				
CMP <sup>1</sup>							
Pregnancy test	Week 4	Week 4	Week 4				
Week 8	HCV RNA	HCV RNA	HCV RNA				
HCV RNA	CBC CMP <sup>1</sup>	CBC CMP <sup>1</sup>	CBC CMP <sup>1</sup>				
CBC	CiviP Pregnancy test	Pregnancy test	Pregnancy test				
CMP <sup>1</sup>	regnancy test	regrandy test	regnutey test				
Pregnancy test	Week 8	Weeks 8 & 12	Weeks 8, 12, 16, & 20				
	CBC	CBC	CBC				
	CMP <sup>1</sup>	CMP <sup>1</sup>	CMP <sup>1</sup>				
	Pregnancy test	Pregnancy test	Pregnancy test				
	Week 12	Week 16	Week 24				
	HCV RNA	HCV RNA	HCV RNA				
	CBC	CBC	CBC				
	CMP <sup>1</sup>	CMP <sup>1</sup>	CMP <sup>1</sup>				
	Pregnancy test	Pregnancy test	Pregnancy test				
Nurse follow-up in clinic	or by phone:						
Managing side effects							
Medication adherence							
Alcohol intake							
Birth control reminde	er						
Refill reminder	3 months post tre	atment 6 months	post treatment				
	CBC		HCV RNA				
	Liver Funct		AFP				
	HCV RNA		RUQ US (if advanced fibrosis)				
	AUDIT-C		AUDIT-C				
	I AUDII-C	<i>F</i>	10DII-C				

<sup>1- &</sup>lt;u>Sofosbuvir- or daclatasvir-based regimen</u> - If GFR <30, no safe recommendation. <u>With ribavirin</u> - If GFR <50, decrease dose (refer to package insert).

Viekira Pak™ (Ombitasvir/Paritaprevir/Ritonavir; Dasabuvir) or Viekira XR ™ (Dasabuvir/Ombitasvir/Paritaprevir/Ritonavir) Treatment Agreement

nily Medicine Provider:
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If you are considering hepatitis C treatment, please read this treatment agreement carefully and be sure to ask any questions you may have before you sign the form.

The FDA approved ombitasvir, paritaprevir, and ritonavir tablets; dasabuvir tablets co-packaged (Viekira Pak™) and dasabuvir, ombitasvir, paritaprevir, and ritonavir extended-release tablets (Viekira XR™) for the treatment of hepatitis C genotype 1b, including those with HCV/HIV co-infection.

Treatment with Viekira Pak™ or Viekira XR™ requires 6 scheduled visits over 6 months for a treatment course of 12 weeks.

#### PREGNANCY & BREASTFEEDING WARNING

It is not known if Viekira Pak™ or Viekira XR™ will harm an unborn or breastfeeding baby, so it is recommended that women do not get pregnant or breastfeed while taking this medicine.

You must stop using ethinyl estradiol-containing medicines before you start treatment with Viekira Pak<sup>™</sup> or Viekira XR<sup>™</sup>. If you use these medicines as a method of birth control you must use another method of birth control during treatment with Viekira Pak<sup>™</sup> or Viekira XR<sup>™</sup>, and for about 2 weeks after finishing treatment with Viekira Pak<sup>™</sup> or Viekira XR<sup>™</sup>. Progestin only contraceptives (e.g. mini pill, Depo shot, Nexplanon<sup>™</sup>) are safe to use during treatment with Viekira Pak<sup>™</sup> or Viekira XR<sup>™</sup>.

#### **HOW THE TREATMENT PROCESS WORKS**

You will have blood and urine tests.

- These tests will include a pregnancy test for female patients. A urine pregnancy test will be done monthly during a clinic visit.
- Random drug and alcohol tests may be requested.
- At each visit, about 2-3 tubes of blood will be collected. Other examinations and tests may be done during the treatment if your provider feels there is a need.

#### Provider, choose appropriate treatment regimen:

Viekira Pak™ or Viekira XR™ will be given for 12 weeks if:

\_\_\_\_\_ You have genotype 1b and do not have cirrhosis.

\_\_\_\_\_ You have genotype 1b and have mild cirrhosis. (Contraindicated in persons with moderate to severe cirrhosis)

Your first visit will be at the start of treatment (week 0) and then visits will be once each month until you stop taking the medications.

You may need to see your primary care provider more frequently if you are having side effects or problems related to the treatment.

You will have a clinic visit 3 months after treatment completion and then yearly (corresponding to your end of treatment date) for 5 years. If you have cirrhosis you should continue to have a liver ultrasound and alpha fetoprotein (AFP) cancer screening blood test every six months and regular clinic visits.

#### TREATMENT MEDICATIONS AND SIDE EFFECTS

<u>Viekira Pak</u><sup>™</sup> contains ombitasvir, paritaprevir, and ritonavir tablets and dasabuvir tablets copackaged for oral use. Two pink tablets contain ombitasvir 12.5mg, paritaprevir 75mg, and ritonavir 50mg and are taken at the same time daily (in the morning) with a meal. Two beige tablets contain dasabuvir 250mg and one of them is taken twice daily (in the morning and evening) with a meal. Store the medication at room temperature.

If you miss a dose of the <u>pink tablets</u>, and it is less than 12 hours from the time you usually take your dose, take the missed dose with a meal as soon as possible. Take your next dose at your regular time the next day with a meal. If you miss a dose of the pink tablets more than 12 hours from the time you usually take your dose, do not take the missed dose. Take your next dose at your regular time the next day with a meal.

If you miss a dose of the <u>beige tablet</u> and it is less than 6 hours from the time you usually take your dose, take the dose with a meal as soon as possible. Then take your next dose at your

regular time with a meal. If it is more than 6 hours since you missed your dose, do not take the missed dose. Instead, take your next dose at your regular time with a meal.

<u>Viekira XR</u><sup>™</sup> contains dasabuvir, ombitasvir, paritaprevir, and ritonavir extended-release tablets, for oral use. Three pale yellow-colored tablets contain dasabuvir 200mg, ombitasvir 8.33mg, paritaprevir 50mg, and ritonavir 33.33mg and are taken at the same time daily with a meal. Taking the tablets in a fasting state could result in reduced cure and the development of resistance. Swallow the tablets whole and do not consume alcohol within 4 hours of taking Viekira XR. Do not skip or miss doses. Store the medication at or below 86°F.

Do not take more than the prescribed dose of Viekira Pak™ or Viekira XR™ (no doubling up). Do not take Viekira Pak™ or Viekira XR™ if you have had a severe skin rash after taking ritonavir (Norvir®).

The most common side effects of Viekira Pak™ or Viekira XR™ are nausea, itching, and sleep problems.

Tell your healthcare provider if you are taking any of the following medicines; as they are contraindicated with Viekira Pak™ or Viekira XR™:

- Alfuzosin hydrochloride (Uroxatral™)
- Colchicine (COLCRYS, Mitigare®)
- Carbamazepine (Carbatrol<sup>™</sup>, Epitol<sup>™</sup>, Equetro<sup>™</sup>, Tegretol<sup>™</sup>); Phenytoin (Dilantin<sup>™</sup>,
   Phenytek<sup>™</sup>); Phenobarbital (Luminal<sup>™</sup>); Primidone (Mysoline<sup>®</sup>)
- Dronedarone (Multag<sup>®</sup>); Lurasidone (Latuda<sup>®</sup>)
- Efavirenz (Atripla<sup>™</sup>, Sustiva<sup>™</sup>)
- Ergot containing medicines including: ergotamine tartrate (Cafergot™, Ergomar™,
   Ergostat™, Medihaler™); dihydroergotamine mesylate (D.H.E. 45™, Migranal™);
   methylergonovine (Methergine®); ergonovine (Ergotrate®)
- Ethinyl estradiol-containing medications; combination birth control pills or patches, such as Lo Loestrin™ FE, Norinyl™, Ortho Tri-Cyclen Lo™, Ortho Evra™; hormonal vaginal rings such as NuvaRing®; hormonal replacement therapy medicine Fem HRT™.
- Gemfibrozil (Lopid™); Lovastatin (Advicor™, Altoprev™, Mevacor™); Simvastatin (Simcor™, Vytorin™, Zocor™)

- Midazolam, when taken by mouth; Triazolam (Halcion™)
- Pimozide (Orap™)
- Rifampin (Rifadin™, Rifamate™, Rifater™, Rimactane™)
- Ranolazine (Ranexa®)
- Sildenafil citrate (Revatio<sup>™</sup>) when taken for pulmonary artery hypertension
- St. John's wort (Hypericum perforatum) or a product that contains St. John's wort

Tell your healthcare provider if you are taking any of the following medicines; as they are <u>not</u> recommended with Viekira Pak™ or Viekira XR™:

- Darunavir (Prezista®) / ritonavir; lopinavir/ ritonavir (Kaletra®); rilpivirine (Edurant®,
   Complera®, Odefsey®)
- Salmeterol (Serevent, Advair®)

Tell your healthcare provider if you are taking any of the following medicines; as <u>dosage</u> adjustments or monitoring may be recommended:

- Amiodarone (Cordarone®, Nexterone®, Pacerone®); bepridil; disopyramide (Norpace®, Norpace CR®); flecainide (Tambocor™); systemic lidocaine (Xylocaine®); mexiletine; propafenone (Rythmol, Rythmol SR); quinidine (Nuedexta®)
- Valsartan (Diovan®, Exforge®, Entresto™); losartan (Cozaar®, Hyzaar®); candesartan
   (Atacand®)
- Amlodipine (Norvasc®); Nifedipine (Procardia®, Adalat®); Diltiazem (Cardizem®,
   Tiazac®); verapamil (Covera-HS®, Calan®, Verelan®)
- Furosemide (Lasix®)
- Ketoconazole; Voriconazole (Vfend®)
- Fluticasone (Inhaled- Arnuity™ Ellipta®, Breo Ellipta®, Flovent®, Advair®; Nasal –
   Flonase®, Veramyst, Dymista®)
- Atazanavir (Reyataz<sup>®</sup>)/ ritonavir
- Cyclosporine (Gengraf<sup>®</sup>, Neoral<sup>®</sup>, Sandimmune<sup>®</sup>); tacrolimus (Astagraf XL<sup>®</sup>, Envarsus XR<sup>™</sup>, FK506 (common name), Hecoria<sup>™</sup>, Prograf<sup>®</sup>)
- Buprenorphine/naloxone (Suboxone®)
- Omeprazole (Prilosec®, Prilosec OTC®)

Alprazolam (Xanax®)

Quetiapine (Seroquel<sup>®</sup>)

Rosuvastatin (Crestor®); pravastatin (Pravachol®)

**PLEASE NOTE:** 

You must let your medical, mental health, dental providers, and pharmacist(s) know that you are taking Viekira Pak™ or Viekira XR™ prior to starting any new medications. You must let your providers know about any new medications you are prescribed before starting them. This

includes vitamins and other supplements.

\*\*\*Hepatitis C treatment should not cause pain that requires narcotic pain medication.

**BENEFITS OF TREATMENT** 

In most cases, hepatitis C will respond to treatment as determined by a blood test that measures the presence and amount of hepatitis C in the blood. If you have no hepatitis C in your blood 12 weeks **after** the end of treatment, this is called a "sustained virologic response" and means you no longer have hepatitis C. Your chance of achieving a sustained virologic response depends on the hepatitis C genotype, how much hepatitis C virus you have in your blood at the beginning of treatment, any past treatment response, and how much liver damage

you have had prior to treatment.

It is possible that you may develop some serious side effects, which will require you to stop the treatment. You may still benefit from treatment even if it does not get rid of your hepatitis C, as

it may slow down the disease. You may choose to stop treatment at any time.

**In Clinical Trials:** 

Persons with genotype 1b, without cirrhosis who were treatment-naïve (never treated before) or treatment-experienced (prior treatment failed), given Viekira Pak™ (the components of Viekira XR™) for 12 weeks had a 100% response (cure) rate. Those with compensated (Child Pugh A) cirrhosis who were either treatment naïve or had prior treatment experience had a 100% response rate.

WHOM TO CALL

If you have any questions about your treatment, contact your primary care provider.

## TREATMENT AGREEMENT

Provider's Name (PLEASE PRINT)	Provider's Signature	Date
Patient's Name (PLEASE PRINT)	Patient's Signature	Date
the information has been explained to	_	and/or the meaning of
My signature below means that I have	·	and/or the meaning of
razors or nail clippers and covering cuts		ecules, toothiblushes,
I will protect myself and others fi	rom hanatitis C by not sharing no	andles toothbrushes
do so, I will contact my provider.	incutions as prescribed by my pre	Triaci. Il I alli ullable tu
I will do my best to take my med		wider If I am unable to
it is in the best interest of my health an	• • •	rider reers triat stopping
I understand that my provider calls		vider feels that stonning
I understand that my hepatitis C	may not respond to treatment	
n i mave any problems with the i provider or nurse know right away.	medications of side effects that	bother me, I will let my
If I have any problems with the i		hother me I will let my
	ally sterile or post-menopausal.	become pregnant.
while on treatment. I understand that n		
rreatment. As a female taking Viekira Pak™ (	or Viokira VP™ Lwill not got prog	mant or broastfood
treatment.	i-being during treatment and the	e effectiveness of
required to evaluate my health and wel	• •	• •
I understand that my treatment		annointments as
this ahead of time and I will reschedule		ice my provider know
length of the treatment. If I am unable	•	
I am willing to visit the clinic and	• • •	•
conditions (depression, history of suicid		
blood pressure, diabetes, high choleste		=
I will tell my provider if I have a		
<b>responses:</b> I agree <u>not</u> to drink alcohol or us	o recreational drugs during the t	roatmont
rosponsos	w the following statements a	and initial beside the

## Viekira Pak™ (Ombitasvir/Paritaprevir/Ritonavir & Dasabuvir) or Viekira XR™ (Ombitasvir/Paritaprevir/Ritonavir/Dasabuvir) Treatment Medications You will take **Viekira Pak™**. Take 2 pink tablets and 1 beige tablet in the morning with a meal. Take 1 beige tablet in the evening with a meal. The generic name for Viekira Pak™ is ombitasvir, paritaprevir, and ritonavir tablets; dasabuvir tablets, co-packaged for oral use. Two pink tablets contain ombitasvir 12.5 mg, paritaprevir 75 mg, and ritonavir 50 mg. The two pink tablets are taken at the same time daily (in the morning) with a meal. The beige tablets contain dasabuvir 250 mg. Take one beige tablet twice daily (in the morning and evening) with a meal. You will take **Viekira XR**™. Take 3 pale yellow-colored tablets each morning with a meal. The generic name for Viekira XR™ is dasabuvir, ombitasvir, paritaprevir, and ritonavir extended release tablets for oral use. These tablets contain dasabuvir 200mg, ombitasvir 8.33mg, paritaprevir 50mg, and ritonavir 33.33mg. You get \_\_\_\_\_\_ from \_\_\_\_\_.

Call \_\_\_\_\_\_ to schedule your family medicine treatment appointments, or if you have any other health concerns.

\*\*\*For any emergencies after normal business hours, please go to the Emergency Room. Make sure any healthcare provider you see knows you are on treatment. Carry a list of your medicines with you.

For more information on managing side effects

visit: http://www.anthctoday.org/community/hep/patients/index.html

Pick up refills on:

Click on "Patient Guide- Managing HepC Treatment"

#### Viekira Pak™ or Viekira XR™ 12 week Lab Tracking Form

#### **General Patient Information**

#### **Pre-Treatment Lab Results**

Med	lication	Regimer

		Select which formulation is being taken:
Name:	HCV RNA:	-Viekira Pak <sup>TM</sup> Do not change dose.  2 pink tablets of ombitasvir, paritaprevir, ritonavir with breakfast.
MRN:	Genotype: HIV: TSH: Vit D 25OH: AFP: GFR:	1 beige tablet of dasabuvir with breakfast and 1 with dinner.  OR
Phone #:	PT/INR: A1C/Glucose:	-Viekira XR <sup>™</sup> Do not change dose.
Treatment Start Date:		3 yellow tablets of dasabuvir, ombitasvir, paritaprevir, ritonavir with a meal.

Completed Treatment											PHQ-9 (Specified	HCV RNA	Weight	Pregnancy
Week	Lab Date	Hgb	Hct	WBC	PLT	ALT	AST	Alk Phos	Total Bili	Creat/GFR	weeks)	(Specified weeks)	(kg)	Test
Pre-Treatment														
Treatment Start Week 0											PHQ-9	HCV RNA		
optional														
optional														
optional														
Week 4												HCV RNA		
optional														
optional														
Week 8														
optional														
optional														
Week 12												HCV RNA		
optional														
3 months post treatment												HCV RNA		

Labs recommended for each follow up visit: CBC, CMP, pregnancy test (females of childbearing age), and HCV RNA as specified.

# Please Remember

Give the End of Treatment Letter to the patient at the completion of treatment.

End of Treatment Letter is found in Treatment Monitoring section on webpage.

12 weeks after treatment completion obtain an <u>HCV RNA</u> to check for a sustained virologic response (SVR). SVR is considered a virologic cure of hepatitis C.

SVR12 Cure Letter is found in Treatment Monitoring section on webpage.

http://anthctoday.org/community/hep/providers/treatment/index.html