

## DRUG AND THERAPEUTICS BULLETIN OF NAVARRE, **SPAIN**

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## SAFE DRUG USE IN PATIENTS WITH CIRRHOSIS

8

Cirrhosis is the fibrosis of the liver as a result of different liver damage mechanisms that leads to the inflammation and generation of scar tissue, thus producing nodules. Liver cirrhosis affects both drug pharmacokinetics and pharmacodynamics, and must therefore be taken into consideration when prescribing. The severity and prognosis of cirrhosis are measured using the Child-Pugh classification.

The aim of this bulletin is to help with selection of the most appropriate therapeutic alternative for patients with cirrhosis by providing a series of safety and dose-adjustment guidelines for the drugs most commonly prescribed in primary healthcare.

When drafting these guidelines, we have based our recommendations on the corresponding summary of product characteristics, UpToDate® and the drug classification based on their safety in liver cirrhosis patients established by a committee of experts from the Netherlands by way of pharmacokinetic studies.

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## index

#### Introduction

Diagnosis and severity of cirrhosis

How does cirrhosis affect pharmacokinetics and pharmacodynamics?

- > Absorption
- > Distribution
- > Metabolism
- > Excretion
- > Pharmacodynamics

#### Can I use this drug in liver patients with cirrhosis?

- > Analgesics
- > Antidepressants
- > Antidiabetics
- > Antihistamines
- > Antimicrobials
- > Antipsychotics
- > Antithrombotics
- > Benzodiazepines and related drugs
- > Systemic corticosteroids
- > Lipid-lowering agents
- > Digestive therapy
- > Cardiovascular therapy

#### Conclusions

Annex 1: Drugs to be avoided in patients with cirrhosis

Annex 2: The situation in Navarre

#### References





#### INTRODUCTION

The liver is the main metabolizer organ in the body and plays a key role in the elimination of numerous substances<sup>1</sup>. Cirrhosis is a chronic and progressive liver disease characterised by fibrosis, alterations of the liver architecture and the formation of regeneration nodules, which occasionally has functional consequences. In advanced stages of cirrhosis, this disease is irreversible<sup>2</sup>. Fibrotic liver diseases in which nodules do not form are not considered to be cirrhosis<sup>1,2</sup>. Liver cirrhosis affects drug metabolism and, as such, it is essential to take this disease into consideration when prescribing<sup>3,4</sup>. Liver failure is the inability of the liver to perform its normal physiological actions, such as the metabolism of endogenous and exogenous substances and the synthesis of plasma proteins<sup>3,4</sup>.

LIVER FAILURE IS A FUNCTIONAL ALTERATION OF THE LIVER, WHEREAS CIRRHOSIS IS A MORPHOLOGICAL ALTERATION OF THIS ORGAN. THERE MAY BE LIVER FAILURE WITH OR WITHOUT CIRRHOSIS AND CIRRHOSIS WITH OR WITHOUT LIVER FAILURE.

There is currently no simple laboratory technique to determine liver function and its severity. Analytical tests do not correlate exactly with liver damage and may even be normal in patients with advanced liver disease. As such, they must be interpreted together with the patient's clinical presentation<sup>2,5,6</sup>.

It is estimated that around 20% of drugs prescribed to patients with cirrhosis are incorrectly dosed or contraindicated and, as a result, 38% of patients

## Liver cirrhosis is characterised by fibrosis and the appearance of nodules in the liver

suffer adverse drug reactions (ADRs), even though approximately 70% of them are preventable<sup>7</sup>. Although several studies concerning pharmacokinetic changes in liver cirrhosis have been published, as yet there is still no guideline with recommendation for the use and dosing of drugs in this situation<sup>8-12</sup>. Moreover, the summary of product characteristics for drugs often do not provide clear management and dosing recommendations, and these recommendations occasionally do not agree with the current safety data<sup>13</sup>.

The aim of this bulletin is to help with selection of the most appropriate therapeutic alternative in liver cirrhosis and to provide a series of safety and dose-adjustment guidelines for the drugs most commonly prescribed in primary healthcare.

#### **DIAGNOSIS AND SEVERITY OF CIRRHOSIS**

There is currently no laboratory test to accurately diagnose liver cirrhosis, and imaging tests are required for its diagnosis. The severity and prognosis are measured using the Child-Pugh classification (Table 1), which gives a view of liver function<sup>14</sup>.

| Parameter  | Score assigned to each parameter |                |            |  |
|--|----------------------------------|----------------|------------|--|
|  | 1                                | 2              | 3          |  |
| Ascites  | Absent                           | Slight         | Moderate   |  |
| Bilirubin (mg/dL)                                  | <2                               | 2-3            | >3         |  |
| Albumin (g/dL)                                     | >3.5                             | 2.8-3.5        | <2.8       |  |
| Prothrombin time<br>Seconds over control or<br>INR | 1-3<br><1.8                      | 4-6<br>1.8-2.3 | >6<br>>2.3 |  |
| Encephalopathy                                     | None                             | Grade 1-2      | Grade 3-4  |  |

#### Table 1. Child-Pugh classification for cirrhosis severity and prognosis.



The total score for the different parameters in the table gives an index, which indicates the degree of liver damage, with higher scores indicating greater severity:

- **5–6 points:** grade A: mild liver failure, well-compensated disease.
- **7–9 points:** grade B: moderate liver failure, with significant functional compromise.
- 10–15 points: grade C: severe liver failure, decompensated disease<sup>14</sup>.

A score of 8 or higher indicates deficient liver function, with potential decompensation and a high risk of complications and death. The Child-Pugh classification is not suitable for estimating the severity of other liver diseases in the absence of cirrhosis<sup>14</sup>.

Patients with well-compensated cirrhosis (Child-Pugh A) do not typically present symptoms, although non-specific symptoms such as fatigue or reduced appetite may appear. The clinical manifestations normally appear in decompensated disease when complications such as ascites present. As such, this disease often goes unnoticed<sup>1</sup>.

#### HOW DOES CIRRHOSIS AFFECT PHARMACOKINETICS AND PHARMACODYNAMICS?

#### Absorption

The portal hypertension that appears during cirrhosis may result in the formation of collateral blood vessels in the liver and portosystemic shunts, thereby increasing the bioavailability of drugs with a high first-pass effect. This results in **elevated plasma concentrations** when these drugs are administered orally<sup>8.15.16</sup>.

#### Distribution

The reduction of albumin and other plasma proteins production in patients with cirrhosis, together with the higher concentration of endogenous substances that attach themselves to the binding sites in albumin, such as bilirubin, causes an **increase in the free fraction** of those drugs that exhibit **high protein binding**. This, in turn, leads to an increase in the effect and/or toxicity, or accelerated elimination, depending on the drug. The presence of ascites may lead to **lower concentrations** of **water-soluble drugs** in systemic circulation as they are distributed into the ascitic fluid<sup>3,8,15,16</sup>. The Child-Pugh classification concerns the severity and prognosis of patients with cirrhosis and is not directly related to liver function

Changes to drug pharmacokinetics and pharmacodynamics occur in liver cirrhosis

#### Metabolism

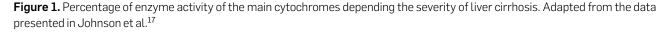
In cirrhosis there is a lower quantity of drug-metabolizing enzymes, although this is not proportional as the cirrhosis worsens and the different cytochromes are also not reduced in the same way<sup>8,15,16</sup>. Figure 1 shows the percentage activity of the main cytochromes involved in drug metabolism according to cirrhosis severity<sup>17</sup>.

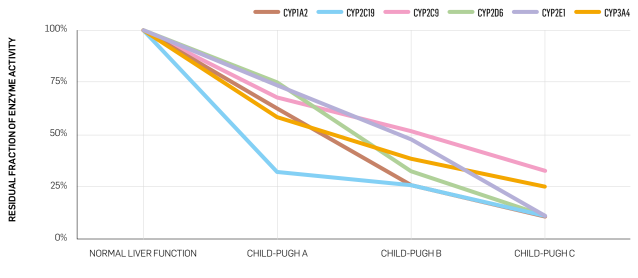
#### Excretion

The excretion of drugs eliminated via the bile duct is reduced, especially in patients with cholestasis. In most cases, this effect is compensated by the kidneys. As such, care must be taken with hepatorenal syndrome<sup>8,15,16</sup>.

#### Pharmacodynamics

Patients with cirrhosis may present differences in the quantity and sensitivity of some receptors, thus altering the effect of some drugs, likediuretics, sedatives or anticoagulants. Moreover, these may be more susceptible to some ADRs, such as nephrotoxicity and clotting abnormalities<sup>8,15,16</sup>.





SEVERITY OF LIVER CIRRHOSIS

## CAN I USE THIS DRUG IN LIVER PATIENTS WITH CIRRHOSIS?

The fact that a drug is hepatotoxic does not always mean that its use is contraindicated in patients with cirrhosis<sup>18</sup>. In such patients, the choice of drug and dose adjustment must be guided by the following factors:

- Indication for drug therapy: optimise the treatment and reduce the number of drugs.
- Pharmacokinetics of the drug.
- · Safety of therapeutic alternatives.
- Comorbidities such as kidney failure, heart failure or alcoholism.
- · Concomitant treatment and risk of drug interactions.
- Interactions with food and medicinal plants, such as grapefruit juice or St. John's wort.
- Severity of the cirrhosis, categorized using Child-Pugh classification.

Once treatment has been established, the onset of signs of decompensation of the cirrhosis and extrahepatic ADRs must be monitored. To that end, the participation of a multi-disciplinary team involving both primary and specialised care professionals is essential to perform dose adjustments and control the effect and tolerance of the treatment.

When preparing this bulletin, we relied on the classification provided by a committee of experts from the Netherlands and published by Weersink et al.<sup>19</sup>, which classifies drugs into six groups on the basis of their safety profile, as based on pharmacokinetic studies in patients with hepatic cirrhosis (Table 2). The recommendations have been checked in the summary of product characteristics<sup>20</sup>, UpToDate<sup>®21</sup> and the database on the Dutch version of the web page *Geneesmiddelen bij levercirrose*<sup>®22</sup>. The groups of drugs classified by the Dutch working group and marketed in Spain at the time this bulletin was drafted have been included. In case of the recommendations differ with the administration route used, the route is specified.

Data concerning the safety of drugs in liver patients with cirrhosis can be consulted at the following sources:



THE RECOMMENDATIONS IN THIS BULLETIN ARE LIMITED TO PATIENTS WITH ESTABLISHED LIVER CIRRHOSIS AND NOT TO OTHER LIVER DISEASES NOT ACCOMPANIED BY CIRRHOSIS.

The drug-induced hepatotoxicity information can be consulted in:



| Classification | Description  | Recommendation  |
|----------------|--|---|
| Safe           | Safety is supported by pharmacokinetic studies and/or long-<br>term safety studies in these patients, with no safety problems<br>having been found.  | Can be used.  |
| No risks known | The limited data suggest that the drug does not increase harm<br>in comparison with patients with no cirrhosis.<br>Drugs classified as "minimally affected by cirrhosis" (with hepa-<br>tic clearance of less than 20%) based on their pharmacokinetics<br>are included. | Can be used.<br>Monitor adverse reactions.<br>Dose adjustment may be required.                            |
| Risks known    | The limited data suggest an increase in patient harm compared<br>with patients with no cirrhosis. However, the number of studies<br>is limited and/or the safety results are contradictory.  | Do not use if a safer alternative is available.<br>If used, possible adverse reactions must be monitored. |
| Unsafe         | The data indicate that this drug is notsafe.   | Avoid.  |
| Unknown        | Insufficient data evaluating the safety are available.   | It should not be used if a safer alternative is available.<br>Evaluate the risk/benefit for each patient. |
| Not classified | The safety has not been evaluated in patients with cirrhosis.  | No recommendations can be made.   |

#### Table 2. Classification of drugs based on their safety in patients with cirrhosis<sup>19</sup>.

#### Analgesics (Table 3)

**Non-steroidal anti-inflammatories (NSAIDs)** must be avoided in patients with cirrhosis regardless of the severity due to the risk of kidney dysfunction and, as a result, decompensation of the cirrhosis. In addition, these patients are at higher risk of suffering gastrointestinal bleeding<sup>22</sup>.

**Paracetamol** is the analgesic treatment of choice. Although its metabolism is slowed in cirrhosis, the hepatotoxic metabolite (N-acetyl benzoquinone imine) does not increase and it can be used safely at non-hepatotoxic doses (up to 4 grammes per day). In patients with riskfactors for hepatotoxicity (such as malnutrition or alcohol consumption), the dose should be limited to 2 g per day<sup>22</sup>. There is currently insufficient evidence available concerning **metamizole**<sup>20,22</sup>.

**Gabapentin** should be started at a low dose and slowly increased depending on the effect and tolerance. Caution must be exercised in patients with hepatic encephalopathy, impaired renal function and patients in concomitant treatment with other central nervous system depressants. Although there are currently no safety data available for **pregabalin** in cirrhosis, it is expected to be safe as hepatic metabolism is minimal<sup>21</sup>.

The first-pass effect and metabolism of the majority of **opioid analgesics** is reduced in patients with cirrhosis, thus increasing the plasma concentrations. As such, the starting dose and/or dosing intervals should be adjusted

## Hepatotoxicity does not always contraindicate drug use in patients with cirrhosis

## Paracetamol is the analgesic of choice

and the dose should be increased gradually and more slowly than in patients without cirrhosis given the clinical effect and ADRs. Moreover, opioids have a greater risk of causing or worsening hepatic encephalopathy in these patients. Codeine and tramadol are prodrugs which are metabolised in the liver and converted into the active metabolite. As tramadol itself is also active, the dose should be adjusted<sup>22</sup>. Another source indicates that tramadol should not be used in Child-Pugh C for the treatment of chronic pain and should be stopped if a worsening of liver function is observed<sup>21</sup>. In the case of codeine, its conversion to morphine decreases markedly in advanced cirrhosis, therefore its use should be avoided<sup>22</sup>.

**Table 3.** Safety and dose adjustment for analgesics in patients with cirrhosis.

|                         |  | CHILD-PUGH A  | CHILD-PUGH B   | CHILD-PUGH C   |  |
|-------------------------|--|---|--|--|--|
| NON-OPIOIDS             |  |   |  |  |  |
| NSAID                   | Safety   |   | Unsafe   |  |  |
|                         | Dose   |   | Do not us  | e  |  |
|                         | Safetyª  |   | No risks kno   | own  |  |
| Gabapentin              | Dose   | Starting in patients with cirrhosis:       Start at ≤ 300 mg per day divided         Standard       into 1–3 doses and increase depending on effect and tolerance         Adjustment if liver function worsens:       Standard  |  |  |  |
| Paracetamol 🖌           | Safety   |   | Safe   |  |  |
| Paracetamol 🖤           | Dose   | Standard  | In patients with hepatotoxicity ri   | sk factors: maximum dose 2 g/day   |  |
| OPIOIDS                 |  |   |  |  |  |
| Dumunananahina          | Safety   |   | No risks kno   | own  |  |
| Buprenorphine           | Dose   | Standard  | Start with ha  | If the dose and increase gradually   |  |
|                         | Safety   | Risks known   |  | Unsafe   |  |
| Codeine                 | Dose   | Standard  |  | Do not use   |  |
|                         | Safety   |   | Safe   | No risks known   |  |
| Fentanyl                | Dose   |   | Start with half the dose and   | increase gradually   |  |
|                         | Safety   |   | Unknown  | 1  |  |
| Hydromorphone           | Dose   | Start with a quarter of<br>the dose and increase No dosing recommendations<br>gradually   |  |  |  |
|                         | Safety   |   | Risks knov   | vn   |  |
| Methadone               | Dose   |   | Standard   | 1  |  |
|                         | Safety   |   | No risks kno   | own  |  |
| Morphine                | Dose (oral)  | Start with half the do  | se and increase gradually  | Start with a quarter of the dose and increase gradually  |  |
|                         | Dose (IV)  | No dose adjustment  | Dou  | uble the dosing interval   |  |
|                         | Safety   |   | No risks kno   | own  |  |
| Oxycodone               | Dose   | Start with half the do  | se and increase gradually  | Start with half the dose and double the interval   |  |
|                         | Safety   | Un  | known  | Unsafe   |  |
| Tapentadol              | Dose   | Standard  | Maximum starting dose<br>50 mg every 8 h (immediate<br>release) or 50 mg every 24<br>h (prolonged release) and<br>increase gradually | Do not use   |  |
|                         | Safety   |   | No risks kno   | own  |  |
|                         |  | Start at 50 mg every 12 h<br>and increase depending<br>on effect and ADR <sup>22</sup>  |  | 25 mg and increase gradually.<br>Im dose: 100 mg every 12 h <sup>22</sup>  |  |
| Tramadol <sup>ь</sup> 🖌 | Dose   | Starting in patients with cirrhosis:<br>ACUTE PAIN:<br>Child-Pugh A: start at 50 mg every 8 h and increase to a maximum of 50 mg per day.<br>Child-Pugh B: start at 25 mg every 8–12 h and increase to a maximum of 100 mg per day divided int<br>Child-Pugh C: Do not use<br>CHRONIC PAIN:<br>Do not use |  | IN:<br>rease to a maximum of 50 mg per day.<br>maximum of 100 mg per day divided into 2–3 doses.<br>o not use<br>AIN:<br>e |  |
|                         | Maintenance after worsening of liver function:<br>Do not use <sup>21</sup> |   |  |  |  |

(a) The classification of gabapentin has been approved by the bulletin's editorial committee.

(b) Tramadol is the weak opioid of choice in cirrhosis.

(✔) Of choice.

### Antidepressants (Table 4)

All selective **serotonin reuptake inhibitors (SSRIs)** are metabolised in the liver and their exposure increases with the severity of cirrhosis. Sertraline and fluoxetine are the most affected. In general, ADRs are dose-dependent, therefore it is recommended to start treatment at a low dose, increasing it gradually depending on the effect and tolerance<sup>21,22</sup>.

The elimination of **venlafaxine** decreases as the severity of cirrhosis increases<sup>22</sup>. **Desvenlafaxine** is considered to be a safer alternative<sup>21</sup>. To date, the safety of other groups of antidepressants has not been evaluated<sup>22</sup>.

|                |        | CHILD-PUGH A  | CHILD-PUGH B   | CHILD-PUGH C  |  |
|----------------|--------|---|----------------|---------------|--|
| Citalonrom .   | Safety |   | No risks known |               |  |
| Citalopram 🗸   | Dose   | Standard  |                | Reduce by 50% |  |
| Escitalopram 🗸 | Safety | No risks known  |                |               |  |
|                | Dose   | Start at 5 mg per day for two weeks then increase. Maximum dose 10 mg per day |                |               |  |
| Fluoxetine     | Safety | No risks known  |                | Unsafe        |  |
| Fluoxeline     | Dose   | Reduce by 50%   |                | Do not use    |  |
| Fluvoxamine 🖌  | Safety |   | No risks known |               |  |
|                | Dose   | Standard Reduce by 50%  |                | Reduce by 50% |  |
| Paroxetine     | Safety | No risks known  |                | Unsafe        |  |
| r ai uxetine   | Dose   | Standard  | Reduce by 50%  | Do not use    |  |
|                | Safety | No risks known  |                | Unsafe        |  |
| Sertraline     | Dose   | Reduce by 50%,<br>maximum dose 100 mg per day.                                | Do not use     |               |  |
| Venlafaxine    | Safety | No risks known  |                | Unsafe        |  |
| ventaraxine    | Dose   | Reduce by 50%   |                | Do not use    |  |
|                | Safety |   | No risks known |               |  |
|                |        | ial dose: 50 mg per day;<br>num dose: 100 mg per day                          |                |               |  |

Table 4. Safety and dose adjustment for antidepressants in patients with cirrhosis.

(a) The classification of desvenlafaxine has been approved by the bulletin's editorial committee.

(✔) Of choice.

#### Antidiabetics (Table 5)

The pharmacokinetics of **insulin** is not affected in patients with liver failure and there is a wide experience of its use in patients with cirrhosis<sup>22</sup>.

**Metformin** can be used safely in Child-Pugh A and B cirrhosis provided the patient does not have other risk factors for lactic acidosis, such as alcohol consumption, dehydration, hypotension, kidney failure or heart failure. In Child-Pugh C metformin levels are considerably increased, therefore the dose should be reduced<sup>22</sup>.

 Table 5. Safety and dose adjustment for antidiabetics in patients with cirrhosis.

|                             |          | CHILD-PUGH A  | CHILD-PUGH B  | CHILD-PUGH C   |
|-----------------------------|----------|---|---|--|
| INSULINS                    |          |   |   |  |
| Insulin                     | Safety   |   | Safe  |  |
|                             | Dose     | Individual, depending   | on blood glucose levels   | s and needs  |
| BIGUANIDES                  |          |   |   |  |
|                             | Safety   | Safe  |   | Risks known  |
| Metformin                   | Dose     | Standard  |   | Reduce by 50%  |
|                             | Comments | Care should be taken  | in patients at risk of lac  | tic acidosis   |
| SULFONYLUREAS               |          |   |   |  |
| Glibenclamide<br>Gliclazide | Safety   | Ν   | lo risks known  |  |
| Glimepiride                 | Dose     | Start at the lowest possible dose and ir  | ncrease gradually depe  | nding on effect and tolerance  |
| MEGLITINIDES                |          |   |   |  |
|                             | Safety   | Unknown   | F   | Risks known  |
| Repaglinide                 | Dose     | Start at 50% of the dose and increase<br>gradually depending on effect and<br>tolerance | Start at 50% of the dose and increase gradually<br>depending on effect and tolerance.<br>Maximum dose: 4 mg a day |  |
| THIAZOLIDINEDIONES          |          |   |   |  |
| Pioglitazone                | Safety   | Ν   | lo risks known  |  |
| riogitazone                 | Dose     | Start at 15 mg once daily and incr  | rease gradually. Maxim  | um dose 45 mg per day  |
| iDPP4                       |          |   |   |  |
| Alogliptin                  | Safety   | No risks known  |   | Unknown  |
| 5 <b>5 1</b>                | Dose     | Standard  |   | No dosing recommendations  |
| Linagliptin                 | Safety   | Ν   | lo risks known  |  |
|                             | Dose     |   | Standard  |  |
| Saxagliptin                 | Safety   | Ν   | lo risks known  |  |
| Canagupan                   | Dose     |   | Standard  |  |
| Sitagliptin                 | Safety   | No risks known  |   | Unknown  |
| onagapan                    | Dose     | Standard  |   | No dosing recommendations  |
| Vildagliptin                | Safety   | Ν   | lo risks known  |  |
| Vitaggiptin                 | Dose     |   | Standard  |  |
| iSGLT2                      |          |   |   |  |
| Canagliflozin               | Safety   | No risks known  |   | Unknown  |
| eanagute2                   | Dose     | Standard  |   | No dosing recommendations  |
|                             | Safety   | Ν   | lo risks known  |  |
| Dapagliflozin               | Dose     | Standard  |   | Start at 5 mg per day and then<br>increase. Maximum dose: 10<br>mg a day |
| Empagliflazin               | Safety   | Ν   | lo risks known  |  |
| Empagliflozin               | Dose     |   | Standard  |  |
|                             |          |   |   |  |

**Sulfonylureas** are metabolised in the liver and bind strongly to plasma proteins. However, extensive experience with their use in patients with cirrhosis confirms their safety<sup>22</sup>. The increased risk of hypoglycaemia with this group of drugs, especially in at-risk patients such as alcohol consumers, severe liver failure or kidney failure, should be taken into consideration.

As regards **meglitinides**, data are only available for repaglinide, the exposure of which increases with the severity of cirrhosis<sup>22</sup>.

The limited data available regarding the use of **pioglitazone** in liver cirrhosis show that there are no differences in terms of exposure and adverse reactions in comparison with patients without cirrhosis<sup>22</sup>.

With regard to **sodium-glucose cotransporter type 2 inhibitors (iSGLT2)**, there are few studies in cirrhosis, although they are well tolerated and it is unlikely that relevant pharmacokinetic changes occur<sup>22</sup>. The risk of urinary infections should be taken into consideration with this group of drugs as they have been associated with decompensation of cirrhosis<sup>21</sup>. To date, there are no safety data for ertugliflozin.

The pharmacokinetic studies published show minimal changes to the pharmacokinetics of **dipeptidyl peptidase 4 inhibitors (iDPP-4)** in liver cirrhosis, although little is known about their long-term safety<sup>22</sup>.

There are currently no data available for **GLP-1 receptor** agonists.

#### Antihistamines (Tabla 6)

There are very few published studies regarding the pharmacokinetic changes of antihistamines in patients with cirrhosis. Except for ebastine and fexofenadine, the remainder are eliminated via the hepatic pathway and therefore their pharmacokinetics are affected. In a study in patients with severe liver cirrhosis, hydroxyzine induced hepatic encephalopathy<sup>22</sup>.

 Table 6. Safety and dose adjustment for antihistamines in patients with cirrhosis.

|                              |        | CHILD-PUGH A                     | CHILD-PUGH B     | CHILD-PUGH C |
|------------------------------|--------|----------------------------------|------------------|--------------|
| Cetirizine<br>Desloratadine  | Safety | No risks known<br>Reduce by half |                  |              |
| Levocetirizine<br>Loratadine | Dose   |                                  |                  |              |
| Cinnarizine                  | Safety | Unknown                          |                  |              |
| Ketotifen<br>Rupatadine      | Dose   | No dosing recommendations        |                  |              |
| Ebastine                     | Safety | No                               | risks known      |              |
| Fexofenadine                 | Dose   | Standard                         |                  |              |
| Hydroxyzine                  | Safety | No risks known                   | Unknow           | 'n           |
| Mizolastine                  | Dose   | Standard                         | No dosing recomm | mendations   |

#### Antimicrobials (Table 7)

With regard to **penicillins**, there are published studies supporting the safety of amoxicillin and amoxicillin/ clavulanic acid in liver cirrhosis. There are no published studies concerning the use of phenoxymethylpenicillin. Cases of neurological ADRs and anomalies in the blood count related to high penicillin doses have been reported in these patients<sup>22</sup>. To date, there are no data on the safety of **cephalosporins**, since they are mainly eliminated renally, liver cirrhosis is not expected to affect their pharmacokinetics<sup>20,22</sup>.

**Macrolides** are mainly eliminated by the liver. However, the pharmacokinetics of azithromycin, clarithromycin

and erythromycin are not significantly affected in cirrhosis. A study reported prolongation of the QT interval with the chronic use of erythromycin in patients with cirrhosis and transjugular intrahepatic portosystemic shunt, although not in patients with cirrhosis and with no shunt. Three studies analysed the safety of clarithromycin in the eradicative treatment of Helicobacter pylori in cirrhosis. The ADRs were mild and similar to those in patients without cirrhosis in all cases<sup>22</sup>.

There are few data available regarding the use of oral **tetracyclins** in these patients, and their use should be avoided<sup>19</sup>. In the case of **quinolones**, several studies support the use of norfloxacin and ciprofloxacin but not levofloxacin<sup>22</sup>.

#### Table 7. Safety and dose adjustment for antibiotics in patients with cirrhosis.

|                                       |        | CHILD-PUGH A              | CHILD-PUGH B             | CHILD-PUGH C  |
|---------------------------------------|--------|---------------------------|--------------------------|---------------|
| PENICILLINS                           |        |                           |                          |               |
| Amoxicillin ±                         | Safety |                           | Safe                     |               |
| clavulanic acid                       | Dose   |                           | Standard                 |               |
| Phenoxymethylpenicillin               | Safety |                           | Unknown                  |               |
| rnenoxymetnytpenicitiin               | Dose   |                           | No dosing recommendation | ons           |
| MACROLIDES                            |        |                           |                          |               |
| Azithromycin                          | Safety |                           | No risks known           |               |
| Clarithromycin<br>Erythromycin        | Dose   |                           | Standard                 |               |
| TETRACYCLINES                         |        |                           |                          |               |
| Doxycycline                           | Safety |                           | Unknown                  |               |
| Minocycline                           | Dose   | No dosing recommendations |                          |               |
| QUINOLONES                            |        |                           |                          |               |
| Ciprofloxacin                         | Safety |                           | Safe                     |               |
|                                       | Dose   | Standard                  |                          |               |
| Levofloxacin                          | Safety | Unknown                   |                          |               |
|                                       | Dose   |                           | Standard                 |               |
| Moxifloxacin                          | Safety |                           | No risks known           |               |
|                                       | Dose   |                           | Standard                 |               |
| Norfloxacin                           | Safety |                           | Safe                     |               |
| Noritoxuoin                           | Dose   |                           | Standard                 |               |
| OTHER                                 |        |                           |                          |               |
| Trimethoprim/                         | Safety |                           | No risks known           |               |
| sulfamethoxazole                      | Dose   |                           | Standard                 |               |
| Fosfomycin (oral)                     | Safety |                           | Safe                     |               |
| · · · · · · · · · · · · · · · · · · · | Dose   |                           | Standard                 |               |
|                                       | Safety |                           | Risks known              |               |
| Metronidazole                         | Dose   | Stand                     | dard                     | Reduce by 50% |
|                                       | 2000   |                           | Limit duration to 2 week | S             |
| Nitrofurantoin                        | Safety |                           | Unknown                  |               |
|                                       | Dose   |                           | No dosing recommendation | ons           |

Although no studies of the pharmacokinetic changes with **fosfomycin** have been performed, it is not expected to cause problems in patients with cirrhosis when administered orally. No safety or dosing conclusion can be reached with the use of **nitrofurantoin**<sup>22</sup>.

**Metronidazole** is extensively metabolised in the liver and its clearance is reduced in patients with cirrhosis, increasing the half-life and risk of accumulation. Longterm treatments increase the risk of encephalopathy, therefore treatment should be limited to a maximum of two weeks<sup>22</sup>.

Various studies supporting the safety of **trimethoprim/ sulfamethoxazole** in cirrhosis have not shown safety problems at low doses (maximum 160/800 mg once daily)<sup>22</sup>.

#### Antipsychotics (Table 8)

There are very few published studies regarding the use of antipsychotics in patients with cirrhosis. These patients are more susceptible to the ADRs for these drugs, including extrapyramidal effects and sedation. The risk of onset or worsening of hepatic encephalopathy is also increased. When used, they should be started at the lowest possible dose and effectiveness and adverse reactions should be monitored; if possible, plasma levels should be measured<sup>22</sup>.

 Table 8. Safety and dose adjustment for antipsychotics in patients with cirrhosis.

|  |         | CHILD-PUGH A   | CHILD-PUGH B                     | CHILD-PUGH C                      |  |
|--|---------|--|----------------------------------|-----------------------------------|--|
| TYPICAL ANTIPSY  | CHOTICS |  |                                  |                                   |  |
| Fluphenazine<br>Flupentixol<br>Perphenazine<br>Periciazine | Safety  |  | Unknown                          |                                   |  |
| Pimozide<br>Tiapride<br>Zuclopenthixol                     | Dose    |  | No dosing recommendations        |                                   |  |
| Haloperidol  | Safety  |  | No risks known                   |                                   |  |
| паторепиот   | Dose    | Start at 50% of the st   | andard dose                      | Start at 25% of the standard dose |  |
| Sulpiride  | Safety  |  | No risks known                   |                                   |  |
| Sulpinde   | Dose    | Si   | tart at the lowest standard dose |                                   |  |
| ATYPICAL ANTIPS  | CHOTICS |  |                                  |                                   |  |
| Amisulpride<br>Cariprazine                                 | Safety  |  | Unknown                          |                                   |  |
| Clozapine  | Dose    |  | No dosing recommendations        |                                   |  |
| Aripiprazol  | Safety  | No risks kno   | own                              | Unknown                           |  |
| Απριριαζοι   | Dose    | Standard   | 1                                | No dosing recommendations         |  |
|  | Safety  |  | Unknown                          |                                   |  |
| Lurasidone   | Dose    | Standard   | Start at 18.5 mg pe              | r day and then increase           |  |
|  | Dose    | Stanuaru   | Maximum dose: 74 mg per day      | Maximum dose: 37 mg per day       |  |
| Olanzapine   | Safety  | No risks kno   | own                              | Unknown                           |  |
| Otalizapine  | Dose    | Start at 5 mg per day and increase depe  | ending on effect and tolerance.  | No dosing recommendations         |  |
| Paliperidone   | Safety  | No risks kno   | own                              | Unknown                           |  |
| Fallperiuone   | Dose    | Standard   |                                  | No dosing recommendations         |  |
|  | Safety  | No risks known   | known                            |                                   |  |
| Quetiapine   | Dose    | Start at 25 mg per day and thenNo dosing recommendationsincrease 25-50 mg per day if required. |                                  |                                   |  |
|  | Safety  | No risks kno   | own                              | Unknown                           |  |
| Risperidone  | Dose    | Start at 0.5 mg twice daily, increasing by   | y 0.5 mg each time if necessary. | No dosing recommendations         |  |

#### Antithrombotics (Table 9)

Haemostasis is altered in patients with cirrhosis, therefore anticoagulants must be used with caution. As the INR increases, determination of the aPTT is not reliable, thus making anticoagulant adjustment more difficult<sup>22</sup>.

Direct oral anticoagulants (DOACs) do not increase the risk of haemorrhage in cases of mild or moderate cirrhosis compared with antivitamin K anticoagulants, heparins or patients without cirrhosis. No extended half-lives or increased exposure to apixaban, dabigatran or edoxaban have been observed in patients with Child-Pugh A or B. However, apixaban and edoxaban are metabolised by CYP enzymes, therefore a cirrhosis-related effect cannot be ruled out and they must be used with caution. In the case of rivaroxaban, exposure in Child-Pugh B patients increases two- to threefold, therefore other, safer alternatives must be selected. There is insufficient information regarding the use of DOACs in Child-Pugh C patients. There is no published information about the use of **fondaparinux**<sup>22</sup>.

With regard to **antiaggregants**, the risk of bleeding and thrombosis must be evaluated on an individual basis. As acetylsalicylic acid is the antiaggregant with the widest use experience, it is the drug of choice. There are currently no data regarding the use of ticlopidine<sup>22</sup>.

Table 9. Safety and dose adjustment for anticoagulants and antiaggregants in patients with cirrhosis.

|                               |        | CHILD-PUGH A   | CHILD-PUGH B                   | CHILD-PUGH C              |
|-------------------------------|--------|--|--------------------------------|---------------------------|
| DIRECT ANTICOAGULANTS         |        |  |                                |                           |
| Apixaban                      | Safety | No risks known   | Ur                             | iknown                    |
| , phasan                      | Dose   | Standard   | No dosing re                   | commendations             |
| Dabigatran                    | Safety | No risks known   |                                | Unknown                   |
| Dabigatian                    | Dose   | Standa   | ard                            | No dosing recommendations |
| Edoxaban                      | Safety | No risks known   | Ur                             | iknown                    |
| Luoxaban                      | Dose   | Standard   | No dosing re                   | commendations             |
| Rivaroxaban                   | Safety | No risks known   | U                              | Insafe                    |
| Rival Oxabali                 | Dose   | Standard   | Do                             | notuse                    |
| ANTIVITAMIN K ANTICOAGU       | JLANTS |  |                                |                           |
| Acenocumarol                  | Safety | No risks known   |                                |                           |
| Kochoodmarot                  | Dose   | Starting dose: 3–4 mg day 1, 2 mg day 2, 1 mg day 3 and monitor INR day 4 $$ |                                |                           |
| LOW MOLECULAR WEIGHT HEPARINS |        |  |                                |                           |
| Dalteparin                    | Safety |  | No risks known                 |                           |
| Enoxaparin                    | Dose   | lf used as trea  | tment, dose twice daily instea | ad of once daily.         |
| Tinzaparin                    | Safety |  | Unknown                        |                           |
|                               | Dose   |  | No dosing recommendations      | 5                         |
| PLATELET ANTIAGGREGAN         | TS     |  |                                |                           |
| Acetylsalicylic acid 🖌        | Safety |  | No risks known                 |                           |
|                               | Dose   | Do not   | t adjust when used as antiagg  | regant.                   |
| Clopidogrel                   | Safety | No risks k   | nown                           | Unsafe                    |
| 0.000.003.01                  | Dose   | Standa   | ard                            | Do not use                |
| Dipiridamol                   | Safety | No risks known   | Known adve                     | erse effects              |
| Dipindumot                    | Dose   |  | Standard                       |                           |
| Prasugrel                     | Safety | Norisks  | known                          | Unknown                   |
| riasugret                     | Dose   | Stand  | ard                            | No dosing recommendations |
| Tipogralor                    | Safety | No risks known   | Ur                             | iknown                    |
| Ticagrelor                    | Dose   | Standard   | No dosing re                   | commendations             |

#### Benzodiazepines and related drugs (Table 10)

Benzodiazepines are metabolized by the liver and their clearance is reduced in cirrhosis. In addition, these drugs increase the risk of hepatic encephalopathy in these patients, especially if they have a history of this condition. As such, their use must be avoided as far as possible. If their use is considered necessary, prioritise the use of lorazepam given its short half-life. It is recommended to start at the lowest possible dose and gradually increase it, closely monitoring ADRs, especially during chronic use<sup>22.</sup>

 Table 10. Safety and dose adjustment for benzodiazepines in patients with cirrhosis.

|                           |        | CHILD-PUGH A                       | CHILD-PUGH B                      | CHILD-PUGH C              |  |  |
|---------------------------|--------|------------------------------------|-----------------------------------|---------------------------|--|--|
| Alprazolam                | Safety |                                    | Risks known                       |                           |  |  |
| Alprazolam                | Dose   | Start at half the                  | standard dose                     | No dosing recommendations |  |  |
| Bromazepam<br>Clorazepate | Safety |                                    | Risks known                       |                           |  |  |
| Flurazepam<br>Loprazolam  | Dose   |                                    | No dosing recommendations         |                           |  |  |
| Brotizolam                | Safety |                                    | Risks known                       |                           |  |  |
| Diotizotam                | Dose   | Start at the lowest available dose | Start at half the                 | standard dose             |  |  |
| Chlordiazepoxide          | Safety |                                    | Risks known                       |                           |  |  |
| Chiordiazepoxide          | Dose   | Start at 1/3 the s                 | tandard dose                      | No dosing recommendations |  |  |
| Clobazam                  | Safety |                                    | Risks known                       |                           |  |  |
| Clobazam                  | Dose   | S                                  | tart at the lowest available dose |                           |  |  |
| Diaman                    | Safety |                                    | Risks known                       |                           |  |  |
| Diazepam                  | Dose   |                                    | Start at half the standard dose   |                           |  |  |
|                           | Safety |                                    | Risks known                       |                           |  |  |
| Lorazepam 🗸               | Dose   | Start at the lowest                | t available dose                  | No dosing recommendations |  |  |
| Lowmotonon                | Safety |                                    | Risks known                       |                           |  |  |
| Lormetazepam              | Dose   | Start at half the                  | standard dose                     | No dosing recommendations |  |  |
| Midazolam                 | Safety |                                    | Risks known                       |                           |  |  |
| Midazotam                 | Dose   | Start at the lowest available dose | Start at half the                 | standard dose             |  |  |
| Zalnidana                 | Safety |                                    | Do not use                        |                           |  |  |
| Zolpidem                  | Dose   |                                    | Do not use                        |                           |  |  |
| 7                         | Safety |                                    | Risks known                       |                           |  |  |
| Zopiclone                 | Dose   | Start at the lowest available dose | Start at half the standard dose   | No dosing recommendations |  |  |

(✔) Of choice.

Both prednisolone and prednisone can be used safely in liver cirrhosis. However, the use of prednisolone should be prioritised as conversion of the prodrug prednisone into prednisolone is reduced in these patients. Serious ADRs and a reduced efficacy have been observed for budesonide, therefore its use should be avoided and other therapeutic alternatives must be used<sup>22.</sup>

Table 11. Safety and dose adjustment for systemic corticosteroids in patients with cirrhosis.

|                |        | CHILD-PUGH A | CHILD-PUGH B | CHILD-PUGH C |
|----------------|--------|--------------|--------------|--------------|
| Budesonide     | Safety |              | Unsafe       |              |
| Budesonide     | Dose   |              | Do not use   |              |
| Prednisolone 🖌 | Safety |              | Safe         |              |
|                | Dose   |              | Standard     |              |
| Drodnicono     | Safety |              | Safe         |              |
| Prednisone     | Dose   |              | Standard     |              |

(🗸) Of choice.

#### Lipid-lowering agents (Table 12)

With the exception of atorvastatin, which has exhibited increased plasma levels and risk of rhabdomyolysis in patients with cirrhosis, the majority of **statins** can be used safely in Child-Pugh A and B even though they are metabolised in the liver. The strongest evidence of safety is with simvastatin<sup>22</sup>. Moreover, simvastatin has been related to clinical benefits such as a lower portal pressure and reduced risk of hepatocarcinoma, thus making it the statin of choice<sup>21,22</sup>. These drugs should be started at the lowest possible dose, with subsequent adjustments based on their effect and tolerance. There are insufficient published data in Child-Pugh C<sup>22</sup>.

There are very few published studies regarding the pharmacokinetic changes of **fibrates** in cirrhosis, despite the

# Simvastatin is the statin of choice

fact they are metabolised in the liver. Gemfibrozil and bezafibrate were studied in a trial and no ADRs were found in Child-Pugh A, although the safety in more severe stages remains unknown. No studies have been performed with fenofibrate<sup>22</sup>.

Exposure to **ezetimibe** increases with the severity of cirrhosis, and it is unsafe in advanced disease<sup>22</sup>.

|               |                    | CHILD-PUGH A   | CHILD-PUGH B  | CHILD-PUGH C   |
|---------------|--------------------|--|---|--|
| STATINS       |                    |  |   |  |
| Atorvastatin  | Safety             |  | Unsafe  |  |
| Allivastatin  | Dose               |  | Do not use  |  |
|               | Safety             | No risk  | s known   | Unknown  |
| Fluvastatin   | Dose               | 0.   | d then increase depending<br>t and ADR  | No dosing recommendations  |
|               | Safety             | No risk  | s known   | Unknown  |
| Pravastatin   | Dose               | 0, ,   | d then increase depending<br>t and ADR  | No dosing recommendations  |
|               | Safety             | No risk  | s known   | Unknown  |
| Rosuvastatin  | Dose <sup>21</sup> | Start at 5 mg per day and then<br>increase depending on effect<br>and ADR  | Start at 5 mg per day and then increase depending on effect and ADR up to a maximum of 10 mg    | Use another alternative; if<br>considered necessary, maximum<br>dose 5 mg  |
|               | Safety             | S  | afe   | Unknown  |
| Simvastatin 🖌 | Dose               | Start at 20 mg per day and then<br>increase depending on effect<br>and ADR | Start at 20 mg per day and then<br>increase depending on effect and<br>ADR. Maximum dose 40 mg. | Start at 20 mg per day and then<br>increase depending on effect and<br>ADR. Maximum dose 40 mg (20<br>mg if decompensated cirrhosis) |
| FIBRATES      |                    |  |   |  |
| Bezafibrate   | Safety             | No risks known   | Unk   | nown   |
| Dezalibiate   | Dose               | Standard   | No dosing rec   | ommendations   |
| Fenofibrate   | Safety             |  | Unknown   |  |
| renombrate    | Dose               |  | No dosing recommendations   |  |
| Gemfibrozil   | Safety             | No risks known   | No risks known Unknown  |  |
| Germibrozit   | Dose               | Standard No dosing recommendations   |   | ommendations   |
| OTHER         |                    |  |   |  |
| Ezetimibe     | Safety             | No risks known   | Uns   | safe   |
|               | Dose               | Standard   | Dono  | bt use   |

#### Table 12. Safety and dose adjustment for lipid-lowering drugs in patients with cirrhosis.

(🗸) Of choice.

#### **Digestive therapy** (Table 13)

There are no published studies regarding the pharmacokinetics and safety of **antacids** in patients with liver cirrhosis. However, given their low absorption and mainly renal elimination, their pharmacokinetics is expected to be relatively unaffected<sup>22</sup>.

The pharmacokinetics of histamine-2 (H2) receptor antagonists is only slightly affected in Child-Pugh A and B patients<sup>22</sup>.

The pharmacokinetics of **some proton-pump inhibitors** (**PPIs**) is markedly affected in patients with cirrhosis due to their extensive hepatic metabolism. The plasma concentrations of pantoprazole and lansoprazole increase between four- and sevenfold. In Child-Pugh C patients,

esomeprazole is the only PPI that has been shown to be safe and is therefore the drug of choice. Omeprazole metabolism is affected by CYP2C10 polymorphisms, whereas esomeprazole metabolism is much less affected. In addition, it is important to ensure the indication for PPI treatment given the greater risk of spontaneous bacterial peritonitis and hepatic encephalopathy for these patients during use. Moreover, the risk of respiratory infections, bone fractures and hypomagnesaemia in long-term treatments with these drugs must be taken into consideration<sup>22</sup>.

As regards **prokinetic** drugs, exposure to metoclopramide increases from Child-Pugh B. The plasma concentrations of domperidone increase threefold in Child-Pugh B and are expected to increase with disease severity, therefore this drug should be avoided<sup>22</sup>.

 Table 13. Safety and dose adjustment for digestive system drugs in patients with cirrhosis.

|  |        | CHILD-PUGH A   | CHILD-PUGH B                   | CHILD-PUGH C          |  |  |
|--|--------|--|--------------------------------|-----------------------|--|--|
| ANTACIDS   |        |  |                                |                       |  |  |
| Magnesium/aluminium hydroxide<br>Magnesium/calcium carbonate | Safety | Unknown  |                                |                       |  |  |
|  | Dose   | No dosing recommendations                              |                                |                       |  |  |
| H2 ANTAGONISTS   |        |  |                                |                       |  |  |
| Famotidine   | Safety | No risks known   |                                |                       |  |  |
| Tamoudine  | Dose   | Standard   |                                |                       |  |  |
| PROTON PUMP INHIBITORS                                       |        |  |                                |                       |  |  |
| Esomeprazole 🖌   | Safety |  |                                |                       |  |  |
|  | Dose   | Standard   |                                | Maximum 20 mg per day |  |  |
| Lansoprazole   | Safety | Unsafe   |                                |                       |  |  |
|  | Dose   | Do not use   |                                |                       |  |  |
| Omeprazole   | Safety | No risks known   |                                | Unsafe                |  |  |
|  | Dose   | Maximum dose: 20 mg per day                            |                                | Do not use            |  |  |
| Pantoprazole   | Safety | Unsafe   |                                |                       |  |  |
| Pantoprazole   | Dose   | Do not use   |                                |                       |  |  |
| Rabeprazole  | Safety | No risks known   |                                | Unsafe                |  |  |
|  | Dose   | Start at 10 mg per day.<br>Maximum dose: 20 mg per day | Maximum dose:<br>10 mg per day | Do not use            |  |  |
| PROKINETIC AGENTS  |        |  |                                |                       |  |  |
| Domperidone  | Safety | No risks known   |                                | Unsafe                |  |  |
|  | Dose   | Standard   | One third of the standard dose | Do not use            |  |  |
| Metoclopramide 🖌   | Safety | No risks known   |                                |                       |  |  |
|  | Dose   | Standard   | 50% of the standard dose       |                       |  |  |

( )Of choice.

#### Cardiovascular therapy (Table 14)

ACE inhibitors and ARA-IIs may increase the risk of hypotension and renal impairment in Child-Pugh B and C patients. Its use is not recommended in patients with cirrhosis and ascites. If used, the initial dose must be as low as possible and blood pressure and kidney function must be closely monitored<sup>22</sup>.

**Calcium channel blockers** are mainly eliminated by the liver and the pharmacokinetics are affected in patients with cirrhosis. Reduced doses are required in these patients, although this is not always possible as the commercial presentations available cannot be fractionated. Cases of cardiogenic shock have been reported in patients with cirrhosis receiving treatment with verapamil and diltiazem after insertion of a transjugular intrahepatic portosystemic shunt. The use of amlodipine, nifedipine and diltiazem should be prioritised as the pharmacokinetics of the former are affected to a lesser extent, adjusting the dose<sup>22</sup>.

The pharmacokinetics of liposoluble **beta-blockers** is markedly affected in liver cirrhosis as they present an extensive first-pass effect and the plasma concentrations may increase. It is recommended to increase the dose gradually, depending on the effect and tolerance<sup>22</sup>.

There is extensive experience with **furosemide** in liver cirrhosis, therefore it is a a safe option. However, ADRs, such as hypokalaemia, hyponatraemia, renal dysfunction, muscle cramps and the onset of hepatic encephalopathy, must be monitored, especially in the first weeks of treatment<sup>22</sup>.

Spironolactone is the **potassium-sparing diuretic** most extensively used in cirrhosis. Exposure to triamterene markedly increases in these patients and cases of severe ADRs, such as megaloblastic anaemia, have been reported, therefore this drug should be avoided<sup>22</sup>.

There are very few published data regarding **thiazide diuretics** and, if used, electrolyte changes must be monitored due to the risk of hepatic encephalopathy<sup>22</sup>.

**Table 14.** Safety and dose adjustment for cardiovascular systemdrugs in patients with cirrhosis.

| CEI  |                          | CHILD-PUGH A  | CHILD-PUGH B   | CHILD-PUGH C  |  |  |
|--|--------------------------|---|--|---|--|--|
| Benazepril<br>Captopril<br>Enalapril                             | Safety                   | No risks known  | Risks known  | Unsafe  |  |  |
| Fosinopril<br>Lisinopril<br>Perindopril<br>Ramipril<br>Quinapril | Dose                     | Start at the lowest pose<br>depending o   | sible dose and increase gradually<br>n effect and tolerance                          | Do not use  |  |  |
| RA-II  |                          |   |  |   |  |  |
| Candesartan  | Safety                   | No risks known  | Risks known  | Unsafe  |  |  |
|  | Dose                     |   | y and then increase gradually  | Do not use  |  |  |
| Eprosartan<br>Irbesartan   | Safety                   | No risks known  | Risks known  | Unsafe  |  |  |
| ii besai tari  | Dose<br>Safety           | Start at the lowest pose  | ible dose and increase gradually<br>Risks known                                      | Do not use<br>Unsafe  |  |  |
| Losartan   | Dose                     |   | ay and then increase gradually   | Do not use  |  |  |
|  | Safety                   | No risks known  | Risks known  | Unsafe  |  |  |
| Olmesartan   | Dose                     | Start at 10 mg per day and then increase gradually  | Start at 10 mg per day<br>and then increase gradually.<br>Maximum dose 20 mg per day | Do not use  |  |  |
| Telmisartan  | Safety                   | No risks known  | Risks known  | Unsafe  |  |  |
| recification   | Dose                     | Start at 20 mg per da<br>Maximum  | iy and then increase gradually.<br>dose 40 mg per day                                | Do not use  |  |  |
|  | Safety                   | No risks known  | Risks known  | Unsafe  |  |  |
| /alsartan  | Dose                     | Start at half the d<br>Maximum  | ose and increase gradually.<br>dose 80 mg per day                                    | Do not use  |  |  |
| ALCIUM CHANNEL BL  | OCKERS                   |   | 31   |   |  |  |
| Amlodipine 🖌   | Safety                   |   | No risks known   |   |  |  |
|  | Dose                     | Standard  | Start at 2.5 mg per day  | Start at 2.5 mg every other day                                 |  |  |
| Barnidipine  | Safety                   |   | Unsafe   |   |  |  |
|  | Dose<br>Safety           |   | Do not use   |   |  |  |
| Diltiazem 🖌  | Dose                     | Start at ha   | No risks known<br>alf the dose and increase gradually depe                           | nding on effect and ADR   |  |  |
|  | Safety                   |   | risks known  | Unsafe  |  |  |
| Felodipine   | Dose                     |   | t 2.5 mg per day   | Do not use  |  |  |
| Lacidipine   | Safety                   |   | Unknown  |   |  |  |
|  | Dose                     |   | Start at 2 mg once per day   |   |  |  |
| Lercanidipine  | Safety                   |   | Unknown  | Unsafe  |  |  |
|  | Dose                     | No dosing   | recommendations  | Do not use  |  |  |
| Nicardipine (oral)   | Safety<br>Dose           |   | Unsafe<br>Do not use   |   |  |  |
|  | Safety                   |   | Da nat use<br>Na risks knawn   |   |  |  |
| Nifedipine 🖌   | Dose                     | Start at ha   | Start at half the standard dose Start at half and double                             |   |  |  |
| Nimodipine (oral)  | Safety<br>Dose<br>Safety |   | Norisks known<br>Start at 30 mg three times per day                                  |   |  |  |
| Nitrendipine   | Dose                     |   | Unsafe<br>Do not use   |   |  |  |
|  | Safety                   | No  | risks known  | Unsafe  |  |  |
| Verapamil (oral)   | Dose                     | Start at 40 mg twice per day and then increase depending<br>on effect and ADR. Maximum interval every 12 hours Do not use |  |   |  |  |
|  | Safety                   | RS Safe   |  |   |  |  |
| Carvedilol 🖌 (m)   | Dose                     | Standard Start at half the standard dose Start at 25% of the standard dose  |  |   |  |  |
| Labetalol (oral) (h)   | Safety                   |   | No risks known   |   |  |  |
|  | Dose                     |   | Start at half the standard do  | ise   |  |  |
| Propranolol ✔ (l)  | Safety<br>Dose           |   | Safe<br>Maximum starting dose: 20 mg 3 tin   | res ner dav   |  |  |
|  | Safety                   |   | No risks known   |   |  |  |
| Sotalol (h)  | Dose                     |   | Standard   |   |  |  |
| ARDIOSELECTIVE BE  |                          |   |  |   |  |  |
| Atenolol 🖌 (h)   | Safety                   |   | Safe   |   |  |  |
|  | Dose<br>Safety           |   | Standard<br>No risks known   |   |  |  |
| Bisoprolol (m)   | Dose                     | Start a   | it 2.5 mg per day <sup>21</sup>  | Start at 2.5 mg per day <sup>21</sup><br>Maximum dose 10 mg/day |  |  |
| Celiprolol (m)   | Safety                   |   | Unknown  |   |  |  |
|  | Dose                     |   | No dosing recommendation   | 15  |  |  |
| Esmolol (h)  | Safety<br>Dose           |   | No risks known<br>Standard   |   |  |  |
|  | Dose<br>Safety           | No  | risks known  | Unsafe  |  |  |
| Metoprolol (l)   | Dose                     | Standard  | Start at 1/3 the standard dose   | Do not use  |  |  |
| hobialal (m)   | Safety                   |   | Unsafe   |   |  |  |
| Nebivolol (m)  | Dose                     |   | Do not use   |   |  |  |
| DOP DIURETICS  |                          |   |  |   |  |  |
| Bumetanide   | Safety                   |   | No risks known   |   |  |  |
|  | Dose<br>Safety           |   | Standard<br>Safe   |   |  |  |
| Furosemide   | Dose                     |   | Standard   |   |  |  |
| DTASSIUM-SPARING   |                          |   |  |   |  |  |
| Amiloride  | Safety                   |   | No risks known   |   |  |  |
| Eplerenone   | Dose                     |   | Standard   |   |  |  |
| Spironolactone 🖌   | Safety                   |   | Safe   |   |  |  |
|  | Dose<br>Safety           | _   | Standard<br>Unsafe   |   |  |  |
| Triamterene  | Dose                     |   | Do not use   |   |  |  |
| HIAZIDE DIURETICS  |                          |   |  |   |  |  |
| Chlortalidone  | Safety                   |   | Unknown  |   |  |  |
| Indapamide   | Dose                     |   | No dosing recommendations  |   |  |  |
| Hydrochlorothiazide  | Safety                   |   | No risks known   |   |  |  |
|  | Dose                     |   | Standard   |   |  |  |
|  |                          |   |  |   |  |  |
|  |                          |   | s of liposoluble beta-blockers is markedly aff                                       |   |  |  |

#### Conclusions

Cirrhosis affects drugs safety as a result of pharmacokinetics and pharmacodynamics changes, and the risk of adverse reactions may therefore increased. Consequently, this must be taken into consideration when prescribing.

In this regard, both prescription medications and over-the-counter drugs and alternative therapies (such as herbal products) that the patient is taking should be considered. It is important to educate patients with cirrhosis that they should consult a healthcare professional before taking any type of medication.

The safety of drugs in liver cirrhosis will depend on the severity of the disease, as estimated using the Child-Pugh classification. As such, it is important that this classification is available in the medical records so that it can be taken into account when prescribing medications. The fact that a drug is hepatotoxic does not always mean that its use is contraindicated in patients with cirrhosis.

Having reviewed the literature, we can state that there are no data regarding the management of the majority of drugs in liver cirrhosis. Available recommendations are occasionally not in agreement in the different sources consulted.

Further studies in this field are therefore required to generate reliable and up-to-date data for the management of patients with liver cirrhosis in a safe way.

## Annex 1 Drugs to be avoided in patients with cirrhosis

A series of drugs that must be avoided in patients with a diagnosis of liver cirrhosis, depending on its severity, and the safest therapeutic alternative, are listed below. It should be noted that the dose of therapeutic alternatives may need to be adjusted depending on the Child-Pugh classification<sup>22</sup>.

| Drug class                  | Avoid   | Child-Pugh | Why is it unsafe?   | Alternative                           |
|-----------------------------|---|------------|---|---------------------------------------|
| NSAID                       | All   | A-C        | Risk of kidney failure<br>and decompensation                                      | Paracetamol                           |
| Weak opioids                | Codeine   | С          | Increased plasma levels   | Tramadol                              |
| Beta-blockers               | Nebivolol   | A-C        |   | Propranolol<br>Atenolol<br>Carvedilol |
|                             | Metoprolol  | С          | Decreased first-pass effect   |                                       |
| Calcium channel blockers    | Barnidipine<br>Nicardipine<br>Felodipine<br>Lercanidipine | A-C        | Pharmacokinetic changes   | Amlodipine<br>Nifedipine<br>Diltiazem |
|                             | Verapamil   | С          |   |                                       |
| Diuretics                   | Triamterene   | A-C        | Increased exposure. Increased risk of megaloblastic anaemia.                      | Spironolactone                        |
| ACEI and ARA II             | All   | B-C        | Risk of hypotension and kidney failure  |                                       |
| Cholesterol-lowering drugs  | Atorvastatin  | A-C        | Increased plasma levels   | Simvastatin                           |
| Antiplatelet agents         | Dipyridamole  | B-C        | Risk of kidney failure  | Acetylsalicylic acid                  |
| Proton pump inhibitors      | Pantoprazole<br>Lansoprazole                              | B-C        | Increased plasma levels due to reduced clearance                                  | Esomeprazole                          |
| Prokinetic agents           | Domperidone   | С          | Increased plasma levels   | Metoclopramide                        |
| Glucocorticoids             | Budesonide  | A-C        | Increased exposure. Lower effect and more severe ADRs in patients with cirrhosis. | Prednisolone                          |
| Benzodiazepines and analogs | All, especially<br>zolpidem                               | A-C        | Risk of hepatic encephalopathy  | Lorazepam                             |
| SSRI antidepressants        | Paroxetine<br>Sertraline                                  | A-C        | Increased exposure. Increase in   | Citalopram<br>Escitalopram            |
|                             | Fluoxetine  | B-C        | dose-dependent ADRs   | Fluvoxamine                           |
|                             | Venlafaxine   | С          |   | Desvenlafaxine                        |

Table adapted from Borgsteede et al<sup>22</sup> ADR: adverse drug reaction.

## Annex 2 **The situation in Navarre**

In June 2023 there were 1326 patients with a diagnosis of liver cirrhosis in Navarre. Of these, 52 were classified as Child-Pugh A (3.9%), 63 as Child-Pugh B (4.8%), 23 as Child-Pugh C (1.7%) and the remainder (89.6%) had no classification in the computerised medical records.

Between April and June 2023, 431 active prescriptions of contraindicated medications in liver cirrhosis were detected in 335 (25.3%) patients. The medicines used are listed in Table 15. A safe therapeutic alternative was available in all cases. Medications contraindicated only in less severe forms of cirrhosis could not be evaluated as the Child-Pugh classification was not available.

| Active substances  | Number of prescriptions  |  |
|--------------------|--|--|
| Pantoprazole       | 144 (33.4%)  |  |
| Lansoprazole       | 22 (5.1%)  |  |
| Total              | 166 (38.5%)  |  |
| Atorvastatin       | 131 (30.4%)  |  |
| Total              | 131 (30.4%)  |  |
| lbuprofen          | 30 (7.0%)  |  |
| Dexketoprofen      | 21 (4.9%)  |  |
| Naproxen           | 11 (2.6%)  |  |
| Condroitin sulfate | 8 (1.9%)   |  |
| Diclofenac         | 8 (1.9%)   |  |
| Etoricoxib         | 5 (1.2%)   |  |
| Celecoxib          | 5 (1.2%)   |  |
| Aceclofenac        | 3 (0.7%)   |  |
| Lornoxicam         | 1 (0.2%)   |  |
| Total              | 83 (21.3%)   |  |
| Zolpidem           | 32 (7.4%)  |  |
| Total              | 32 (7.4%)  |  |
| Nebivolol          | 4 (0.9%)   |  |
| Barnidipino        | 4 (0.9%)   |  |
| Nitrendipino       | 2 (0.5%)   |  |
| Total              | 10 (2.3%)  |  |
|                    | Pantoprazole<br>Lansoprazole<br><b>Total</b><br>Atorvastatin<br><b>Total</b><br>Ibuprofen<br>Dexketoprofen<br>Dexketoprofen<br>Condroitin sulfate<br>Diclofenac<br>Diclofenac<br>Colecoxib<br>Aceclofenac<br>Colecoxib<br>Aceclofenac<br>Colecoxib<br>Aceclofenac<br>Colecoxib<br>Aceclofenac<br>Colecoxib<br>Aceclofenac<br>Colecoxib<br>Aceclofenac<br>Colecoxib<br>Aceclofenac<br>Colecoxib<br>Aceclofenac<br>Colecoxib<br>Aceclofenac<br>Colecoxib |  |

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