Prioritization of Therapeutic Targets and Trial Design in Cirrhotic Portal Hypertension


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Abstract

Portal hypertension is the main driver of cirrhosis decompensation, the main determinant of death in patients with cirrhosis. Portal hypertension results initially from increased intrahepatic vascular resistance. Later, increased inflow from splanchnic vasodilation and increased cardiac output lead to a further increase in portal pressure. Reducing portal pressure in cirrhosis results in better outcomes. Removing the cause of cirrhosis might improve portal pressure. However, this is a slow process and patients may continue to be at risk of decompensation. Additionally, for some chronic liver diseases, such as non-alcoholic fatty liver disease, etiological treatments are not yet available. Therefore, there is a need to develop better therapies specifically aimed at reducing portal pressure. For over 35 years, the mainstay of such therapy has been the use of nonselective beta-blockers that act by reducing portal venous inflow. Recently, many drugs (mainly targeting intrahepatic mechanisms) have shown promise in pre-clinical and early clinical studies and may act alone or synergistically with nonselective beta-blockers in reducing portal pressure in cirrhosis. The objective of this position paper is to propose a novel framework for the design of clinical trials (phase 1, 2 and 3) in patients with cirrhosis and portal hypertension and to prioritize novel targets and pharmacological therapies in this setting. We have focused the discussion on patients with
compensated cirrhosis. The paper summarizes discussions held at The American Association for the Study of Liver Diseases (AASLD) Industry Colloquium in January 2018, with the participation of clinical and translational investigators, regulatory professionals and industry partners.

I. Introduction

Portal hypertension (PH) is the main consequence of cirrhosis and is the main driver of cirrhosis decompensation (1). Cirrhosis decompensation is defined by the presence of clinical complications (ascites, variceal hemorrhage, encephalopathy) and constitutes the main determinant of death in patients with cirrhosis (2).

PH results initially from increased intrahepatic resistance due to architectural distortion and intrahepatic vasoconstriction (3). At this stage, PH is mild but leads to increase in vasodilators in the splanchnic vasculature that result in increased portal venous inflow. This increased inflow leads to a further increase in portosystemic gradient and, once it surpasses a certain level (clinically significant portal hypertension), the patient becomes more susceptible to decompensation (4).

Elimination of the underlying cause of cirrhosis may lead to a reduction in fibrosis and to an improvement in PH. For example, elimination of the hepatitis C virus has been associated with a reduction in portal pressure in patients with cirrhosis (5, 6). However, once PH becomes clinically significant, it is unlikely to regress, at least in the short term, and the patient continues to be at risk of decompensation (6, 7). Additionally, there are etiologies (e.g. non-alcoholic fatty liver disease) for which etiological treatment is not yet available.

Reducing portal pressure in cirrhosis, independent of elimination of the cause of liver damage, results in better outcomes (8, 9). Therefore, there is still a need to develop therapies that reduce portal pressure by targeting its main pathogenic vascular mechanisms. For over 35 years, the mainstay of such therapy has been the use of nonselective beta-blockers (NSBB) that act by reducing portal venous inflow.

Recently, many drugs (mainly targeting intrahepatic mechanisms) have shown promise in pre-clinical and early clinical studies. The American Association for the Study of Liver Diseases (AASLD), at its yearly Industry Colloquium (January 19-20, 2018), invited members of the AASLD Portal Hypertension Special Interest Group (SIG) to participate in a symposium that had the objective of discussing novel therapies in PH from the perspective of clinical and translational investigators, with participation from regulatory professionals and industry partners.

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The objective of this position paper, as discussed in the symposium, is to propose a novel framework for the design of clinical trials (phase 1, 2 and 3) in patients with cirrhosis and PH and to prioritize novel targets and pharmacological therapies for PH.

II. Preclinical studies in portal hypertension

The initial mechanism in the pathogenesis of PH, an increase in intrahepatic vascular resistance, is due to a combination of structural changes (i.e., fibrosis, disruption of sinusoidal structure, microthrombi) and increased sinusoidal resistance (i.e. vasoconstriction) that results from increased contraction of activated hepatic stellate cells in the setting of increased vasoconstrictors and to dysfunctional endothelial cells with decreased production of vasodilators such as nitric oxide (NO) (3). In response to intrahepatic changes and/or mild increases in portal pressure, splanchnic and systemic vasodilation and angiogenesis follow, leading to increased portal venous inflow and worsening of PH (10, 11).

The complexity of PH is evident by the divergent actions of vasoconstrictive and vasodilating systems in the intrahepatic vs. the extrahepatic circulations. Vasoconstrictors such as angiotensin II and norepinephrine contribute to intrahepatic vasoconstriction (10). Rho-kinase, a key pathway in the maintenance of basal vascular tone, is dysregulated in both the intrahepatic and the extrahepatic circulations; while in the liver it contributes to vasoconstriction, in the extrahepatic vessels contractility is almost completely lost leading to vasodilatation (10). Conversely, availability of the vasodilator NO is reduced in the liver and increased in the extrahepatic tissues (3). Due to these and other changes, even the strongest vasoconstrictors like angiotensin II or epinephrine fail to constrict extrahepatic vessels, while the intrahepatic vascular bed shows exaggerated contraction. Therefore, it becomes a major challenge to select systemic therapies for PH in cirrhosis because vasoconstrictors (that would target splanchnic vasodilatation) will worsen intrahepatic resistance and vasodilators (that would target intrahepatic vasoconstriction) will lead to systemic hypotension and its downstream deleterious consequences.

Systemic and splanchnic vasodilatation is a key component of the hyperdynamic syndrome in cirrhosis and is characterized clinically by low blood pressure and increased cardiac output. It not only leads to the development of clinically significant PH and portosystemic collaterals but is also responsible for sodium and water retention, all of which eventually lead to the development of clinical decompensation (12). In a stage of further progression of liver disease, systemic hypotension develops mainly due to further vasodilatation and a relative decrease in cardiac output, leading to
refractory ascites and hepatorenal syndrome (13). Systemic hypotension on the one side and clinical immunological events (e.g. systemic inflammation, infections) on the other side lead to impaired organ perfusion and dysfunction, and the development of multiorgan failure in cirrhosis (the so-called acute-on-chronic liver failure) (14). Therefore, the selection of targets could be different in compensated and decompensated cirrhosis depending on the objective of treatment (Figure 1).

Table 1 provides a list of drugs that, in pre-clinical studies, have been shown to reduce portal pressure, including their purported site of action and other potential benefits as well as potential deleterious effects. Of them, only statins have gone on to proof-of-concept clinical trials leading to a randomized controlled trial (RCT) with clinical outcomes (15-17). Statins, are approved drugs for the treatment of hypercholesterolemia but they are pleiotropic drugs that are also Rho-kinase inhibitors and have anti-fibrotic effects. The use of statins is an example of how drugs that are in clinical use for other indications can be repurposed for the treatment of PH and this constitutes a promising model.

The complexity of the processes involved in PH and in the development of complications of cirrhosis would favor the pursuit of multi-targeted therapies such as statins or of combination therapies addressing different targets. An attractive possibility is cell-specific targeting of important pathways. Further research and development is still needed but there are currently several avenues to pursue (table 1).

III. Pharmacokinetic (PK) and pharmacodynamic (PD) evaluation in portal hypertension clinical trials (Phase 1 and Phase 2)

PK studies are generally undertaken during Phase1b and/or Phase2a clinical trials. In addition to investigating PK in healthy volunteers, there should be close attention to understanding the compound’s pharmacokinetic characteristics in hepatically impaired populations. Several pathophysiological changes which regularly occur in individuals with cirrhosis and PH can significantly impact drug disposition in this population (Table 2) (18-23). The 2003 Food and Drug Administration (FDA) Guidance for Industry (24) recommends a formal PK study in hepatically impaired individuals if (a) hepatic metabolism and/or excretion accounts for > 20% of the absorbed drug’s or its active metabolite’s disposition, or (b) if the test compound is either known or suspected to have a narrow therapeutic window, or (c) if there is insufficient information about the compound’s metabolism and excretion and its predicted therapeutic window. However, a formal PK study in hepatically impaired may not be necessary if a compound is entirely renally excreted with
no involvement of the liver, or if a drug is gaseous or volatile and its active metabolite(s) are primarily eliminated via the lungs (24).

PK studies can have a basic full study design, a reduced study design, or a population PK approach. The basic full study design involves the entire Child-Pugh spectrum (at least 6 patients in each Child-Pugh category) and controls, whereas a reduced study design allows for the study to be limited to eight patients with Child-Pugh B cirrhosis and eight controls matched by age, gender, and co-morbidities. The findings from a reduced study design conducted on Child-Pugh B patients, while considered applicable to individuals with Child-Pugh A cirrhosis, would be generally contraindicated in Child-Pugh C patients. Although widely employed, limiting PK studies to a reduced study design is not recommendable as individuals with Child-Pugh A (mostly compensated) and Child B (mostly decompensated) cirrhosis may have markedly different pathophysiological abnormalities (e.g., plasma protein binding, hepatic blood flow, or intrinsic clearance). One might consider a staged approach to a basic full study design where individuals with Child-Pugh A and B are investigated initially, assuming that initial efficacy and safety studies are limited to patients with mild to moderate Child-Pugh cirrhosis. Conducting PK studies in individuals with severe liver dysfunction is challenging due to their tenuous health status with frequent complications and hospitalizations and multiple concomitant medications. Wherever possible, controls in these studies should not be healthy volunteers but should be matched for age, gender, co-morbidities and concomitant medications. A single-dose PK study might be sufficient if a compound and its metabolite exhibits linear and time-independent PK at concentrations anticipated in the patients to be studied and are not expected to change in patients with cirrhosis and PH. Although their validity is not fully established in patients with cirrhosis and PH, population pharmacokinetic analyses should be built into Phase3 and Phase4 studies wherever possible as such data can be quite instructive in fully understanding the PK characteristics.

Patients with cirrhosis may respond differently than the general population to medications due to accompanying pathophysiological abnormalities. These include plasma protein binding, decreased skeletal muscle mass, alterations in blood brain barrier, reduced renal blood flow, and increased QT interval. Some examples of altered PD in cirrhosis include: unresponsiveness to loop diuretics in patients with refractory ascites (25), risk of acute kidney injury from non-steroidal anti-inflammatory drugs in patients with cirrhosis especially ascites (26), and more recent description of worsened insulin sensitivity from nadolol in patients with cirrhosis (27). PD assessments should be considered on a compound by compound basis, especially if concentration-response relationship is unknown, or if altered hepatic function and/or PH are suspected to alter PD response. It is our understanding that the FDA is in the midst of revising its guidance for PK studies in hepatic impaired population. It is
possible that MELD score (rather than Child-Pugh classes) forms the basis for future PK studies in patients with cirrhosis.

**Safety Monitoring in Cirrhosis and Portal Hypertension Clinical Trials**

Monitoring for hepatic and extrahepatic toxicity is very important in drug development for cirrhosis and PH. Two important considerations in monitoring for hepatotoxicity are (a) challenges in biochemical monitoring as baseline aminotransferases and bilirubin may be abnormal and (b) due to lower functional hepatic reserve, patients may be at risk for rapid decompensation and death, even with injuries that are otherwise not as serious (for example, cholestatic liver injury) (28). Patients with cirrhosis may be at risk for extrahepatic (off target) adverse events due to systemic pathological changes such as decreased plasma protein, altered volume of distribution, decreased skeletal muscle mass, decreased renal blood flow, and increased blood brain permeability, and altered gut permeability. Further, these patients may already be taking other medications and thus may be exposed to drug-drug interactions. Every study should have a well thought out plan for safety monitoring and stopping rules and the data safety and monitoring boards should include hepatologists with familiarity in drug safety and causality adjudication.

IV. Proof-of-concept studies (phase 2) trials in portal hypertension

a) Selection of relevant population

Cirrhosis is subclassified in two main clinical stages, compensated and decompensated, each of them with different prognosis and different predominant pathogenic mechanisms and therefore different therapeutic targets (29). Patients with compensated and decompensated cirrhosis should be investigated or analyzed separately in clinical trials.

Compensated cirrhosis is the asymptomatic and the longest stage of cirrhosis. In this stage, the patient with chronic liver disease and/or chronic risk factors for liver disease (e.g. alcohol ingestion, HCV infection) has histological, imaging or elastographic evidence of cirrhosis but has not developed any decompensating events (ascites, variceal hemorrhage or encephalopathy). Median survival is around 15 years (30).

Compensated cirrhosis is sub-classified by the severity of PH into those with mild portal hypertension (MPH) and those with “clinically significant” portal hypertension (CSPH). These two stages are different not only in the degree of PH but also regarding their risk of progression to
decompensation, the amount of hepatic fibrosis and the predominant pathogenic mechanisms. While patients with MPH have a very low risk of decompensation, have less liver fibrosis (and thinner fibrous septa) and the main pathogenic mechanism is increased intrahepatic resistance, those with CSPH have a 4-times higher risk of decompensation than patients with mild PH (1), have more liver fibrous tissue (31) and, in addition to intrahepatic resistance, a main contributor to PH is increased splanchnic blood flow (4). Because non-selective beta-blockers (NSBB) act by decreasing flow, the portal pressure response to NSBB is much lower in patients with MPH compared to those with CSPH (4).

Treatment of underlying disease, if possible, is the most effective and appropriate treatment at the compensated stage. In patients with mild PH, it is conceivable that this treatment (or antifibrotic treatment when available) could lead to regression to a non-cirrhotic stage or at least to halting the progression to CSPH. However, although treatment of the primary liver disease may slow progression to decompensation in patients with CSPH, the risk is not eliminated and a substantial number of patients remain with CSPH and at risk for decompensation. It is in these patients and in those for whom specific etiologic therapies are unavailable where measures to reduce portal pressure will be more important.

**Decompensated cirrhosis** is the symptomatic and shorter stage of cirrhosis. This stage is characterized by the presence of clinical complications, specifically ascites, variceal hemorrhage, encephalopathy and jaundice (although the latter is very rare as first decompensating event) (29). Median survival is 1.5 years (30). The type and number of decompensating events is of prognostic importance as is the presence of further complications of decompensating events such as hepatorenal syndrome (“further” decompensation”). Decompensated patients have, by definition, CSPH. Although worsening vasodilatation and inflammation contribute to outcomes in these patients, PH still plays a major role as outcomes improve with reduction in portal pressure (8, 32).

**b) Measurement of portal pressure and definition of mild vs. CSPH**

The hepatic venous pressure gradient (HVPG) is the main method used to measure portal pressure. It consists of the percutaneous catheterization of a hepatic vein (mostly via the transjugular route) with measurements of the wedged hepatic vein pressure (WHVP) and the free hepatic vein pressure (FHVP) (33). The WHVP is obtained by either advancing the catheter until it is wedged in the most distal portion of the hepatic vein or, preferably, by inflating a balloon at the end of a catheter in a more proximal portion of the vein. It is a measure of hepatic sinusoidal pressure and correlates closely with portal pressure in viral, alcoholic and non-alcoholic steatohepatitis cirrhosis where there

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is sinusoidal and/or post-sinusoidal fibrosis. The FHVP is a measure of systemic pressure and is used as an internal zero to “correct” the WHVP. The HVPG is therefore obtained by subtracting the FHVP from the WHVP. Normal HVPG is 3-5 mmHg. An HVPG > 5 mmHg indicates the presence of PH and an HVPG ≥10 mmHg indicates the presence of CSPH.

Although all patients with cirrhosis should theoretically have an abnormal portal pressure (i.e. HVPG > 5 mmHg), the HVPG is normal in ~10% of patients with histological/clinical cirrhosis. This could be due to sampling error in the biopsy, a predominantly presinusoidal fibrotic component, or to an inadequate occlusion of the hepatic vein due to the presence of intrahepatic shunts.

Sub-classification of patients into mild vs. CSPH is based on the HVPG. Mild PH is defined as an HVPG >5 mmHg but less than 10 mmHg. CSPH is defined as an HVPG ≥ 10 mmHg. Because patients with varices and/or decompensation have an HVPG of at least 10-12 mmHg, patients with varices (or collaterals on imaging) and/or clinical decompensation by definition have CSPH (12).

Non-invasive approaches to assess the presence of CSPH include measurements of liver stiffness with or without additional parameters such as platelet count and/or spleen size (34). The presence of varices that have not bled or a history of variceal hemorrhage can be used to select candidates for studies of patients with CSPH. However, standard of care would require that some of these patients (those with moderate/large varices) receive therapies to prevent variceal hemorrhage (either NSBB and/or ligation) that could confound the interpretation of results. If included in a single study, patients with compensated and decompensated cirrhosis would require separate analysis.

c) Definition of HVPG response

Drugs identified in pre-clinical studies with a portal pressure-reducing effect (candidate drugs) should be tested on their ability to reduce HVPG in patients with CSPH. Unfortunately, until now, there is not a single non-invasive test (e.g. Doppler, liver stiffness) whose changes correlate with changes in HVPG and that would be acceptable alternatives to paired HVPG measurements.

The magnitude of change in the HVPG required to predict that a drug has potential clinical utility in patients with CSPH is still controversial. In patients with a history of variceal hemorrhage, a decrease in HVPG to less than 12 mmHg or a decrease greater than 20% from baseline is associated with a significant reduction in the risk of recurrent hemorrhage, ascites, encephalopathy and death (8). Reductions in HVPG >10% from baseline have been associated with a reduction in variceal development (35), first variceal hemorrhage and death (9).
It is important to note that these threshold HVPG reductions apply to NSBB-based therapies, which act by decreasing portal blood inflow and may not apply to drugs that decrease intrahepatic resistance which, while decreasing portal pressure only modestly may have a mild effect in preventing bleeding, but by improving liver perfusion may have a more important effect on survival (16, 17).

Importantly, using the most common cutoffs (reduction of HVPG to <12 mmHg or >20% from baseline) to classify patients in a dichotomous fashion into HVPG “responders” and “non-responders” may underestimate the value of a candidate drug. While “responders” have in fact been shown to have better clinical outcomes than non-responders, the definition lacks sensitivity as many patients not meeting these criteria may still have a good clinical response to therapy (36). It would be unwise to rule out a potential therapeutic benefit for a drug that achieves a consistent decrease in portal pressure, even if this decrease in portal pressure does not reach the threshold used to define a good response.

On the other hand, preliminary analyses of a dynamic HVPG model obtained from patients with compensated cirrhosis without varices included in the timolol study, demonstrates that, at any time during follow-up, a 1 mmHg increase (or decrease) in HVPG is associated with a 1.19-times higher (or lower) risk of decompensation/death (95%CI:1.11-1.27) (37). This model provides much more granularity to changes in HVPG and, if confirmed, a proof-of-concept study could be powered on a pre-defined specific reduction in HVPG associated with a certain percentage reduction in decompensation in the patient population to be analyzed. Furthermore, if the proof-of-concept study includes clinical outcomes, results could inform a potential effect size for a phase 3 trial.

**d) Comparator to be used**

In proof-of-concept studies that investigate the effect of a candidate drug on HVPG, the use of a placebo of the candidate drug(s) is mandatory.

If the investigational drug has a different mechanism of action from the standard of care (NSBB in patients with medium/large varices), patients on these drugs should be stratified separately to investigate the presence of a synergistic or additive effect of the candidate drug.
e) **Readouts other than HVPG to be investigated in proof-of-concept (POC) studies**

(Table 3)

Candidate drugs with a purported intrahepatic mechanism of action (i.e. reduction of intrahepatic resistance) may have a more important effect by improving liver perfusion while demonstrating an only modest effect on portal pressure. Therefore, analysis of these drugs should include quantitative tests of liver perfusion such as the indocyanine green clearance test, or breath tests that analyze clearance of ingested or injected markers.

Other readouts that should be prospectively collected are the development of clinical outcomes during the course of the study (clinically overt decompensating events) as well as potentially negative readouts such as reduction in mean arterial pressure or cardiac output or development of renal dysfunction, and other adverse events.

In addition, analyses that would support the purported mechanism of action of the candidate drug would be desirable as a correlation between the hemodynamic effect and mechanistic markers would support advancing to a phase 3 trial. These analyses may disclose useful biomarkers of the response to this specific drug.

As in any clinical trial evidence of hepatotoxicity or other adverse events should be collected in the knowledge that the study population (compensated cirrhosis) may have low tolerance to even mild adverse events.

f) **Other considerations**

- **Duration of therapy:** Depends on the proposed mechanism of action of the candidate drug. If it has direct vasoactive action, the HVPG response could be assessed acutely after a short period as has been demonstrated after the intravenous administration of propranolol (9). However, it would be necessary to continue the drug and have a follow-up HVPG after longer administration to demonstrate that the effect is maintained and is not offset by compensatory mechanisms (as occurs with long-acting nitrates). Drugs targeting the structural component of the increased hepatic resistance (e.g. antifibrotic drugs) may take much longer (e.g., 12 months) to demonstrate HVPG reductions.

- Confounders such as concomitant treatment of underlying cause of liver disease (e.g. weight loss, antiviral therapy), co-morbidities (e.g. alcohol use, cardiovascular
disease) or treatment of co-morbidities that may affect HVPG (e.g. statins, vasoactive medications) should be considered.

In summary, POC studies in PH should include patients with cirrhosis and CSPH, with the main readout being a decrease in HVPG. Other readouts (positive and negative) and findings should inform the decision to move on to phase 3 trials.

**V. Phase 3 trials in portal hypertension**

Most clinical trials in PH have been directed towards treating or preventing variceal hemorrhage. However, varices and variceal hemorrhage are not isolated events and they must be considered in the context of the presence (or absence) of other decompensating events (2, 12). Therefore, the aim of clinical trials in PH should be to prevent, not only variceal hemorrhage, but any form of decompensation. Unfortunately, a paucity of data currently exists in this area, and RCTs are needed. Trial design in patients with decompensated cirrhosis has been extensively discussed elsewhere (2) and we therefore focus the discussion on patients with compensated cirrhosis.

**a) Primary endpoint**

In agreement with the consensus achieved at the 2015 Baveno workshop on PH (2), the development of a decompensating event (ascites, variceal hemorrhage and/or encephalopathy) should be the main primary endpoint in Phase 3 trials. Such composite endpoint provides certain advantages. Most importantly, the combination of clinical variables provides the most clinically relevant measure of clinical well-being. Additionally, in RCTS, a composite endpoint will enhance the ability to detect differences in outcomes. It should be noted that ascites is the most common first decompensating event (followed by variceal hemorrhage and encephalopathy (38)). An objective and standardized definition of all these events is needed to adequately assess the main endpoint. Ascites, or free fluid in the peritoneal cavity, should be documented by imaging techniques and confirmed by diagnostic paracentesis (which is otherwise indicated in patients first presenting with ascites), as clinical signs of ascites are not accurate (39). Variceal hemorrhage should follow the definitions of the Baveno VI consensus conference (2). “Covert” hepatic encephalopathy is difficult to detect outside of specialized dedicated units, and reproducibility of this diagnosis is low (40). Therefore, encephalopathy should be “overt” (at a minimum confusion and asterixis, or West Haven Grade ≥2) and documented by a physician to be considered an endpoint. Many patients with
compensated cirrhosis are taking sedatives, anxiolytics or antidepressant drugs that may interfere with the recognition of mild encephalopathy.

b) Development of varices as a part of the main primary endpoint

Although the development of varices without decompensation is a sign of disease progression, this clinical feature should not be included as part of the primary endpoint. Even though the 2-year risk of decompensation in patients with and without varices is different (~15% vs. ~8%, respectively) (1, 30, 38, 41), there are several operational reasons for not considering varices as part of the primary endpoint. First, the trial would be limited to patients without varices, which have a much lower risk of decompensation than patients with varices. Second, the inter-observer agreement for small varices is low (42). Third, if development of varices is a component of a composite primary endpoint, development of varices would lead to the discontinuation of the study drug. This could result in missed opportunities when evaluating the efficacy of drugs that might prevent decompensation, but not the development of varices. In fact, in the study comparing timolol or placebo for the prevention of varices (35), the development of varices led to drug discontinuation, precluding assessment of a potential effect on the development of ascites or other decompensating events. Very recent data, presented in abstract form, suggests that NSBB might have a strong effect preventing ascites in patients with cirrhosis and CSPH (43).

The development of varices should be assessed as a secondary endpoint, but should not be part of the primary endpoint and should not lead to study drug discontinuation. This secondary endpoint could be analyzed as an exploratory ordinal outcome (considering death as the worst potential outcome, followed by decompensation and varices (Fig 2). Patients with varices fulfilling current recommendations for treatment either at baseline or during the trial should receive standard therapy to prevent first variceal hemorrhage, which in the context of a phase 3 drug therapy-based trial should preferably be endoscopic therapy. Ideally this should be standardized across centres.

c) Use of a surrogate endpoint

Using an outcome surrogate is attractive for studies where a low event rate is expected within the study timeframe. A surrogate would not only reduce sample size and study duration but could also be considered appealing because it would be a substitute for a more severe clinical outcome. Because patients with compensated cirrhosis have an overall low rate of decompensation (30, 38), it
has been suggested that changes in HVPG could be used as a surrogate endpoint to support the approval of new treatments for PH (within the FDA accelerated approval framework) (44)).

Although, as mentioned previously, demonstrating a decrease in HVPG is the main goal of phase II trials, several issues would need to be resolved before a drug could be approved based on HVPG response. First, the magnitude of HVPG response considered clinically relevant remains to be determined. Second, most of the data demonstrating an association between an HVPG decrease and a reduction in clinical outcomes were obtained in studies using NSBB (8), drugs that decrease portal pressure by decreasing portal blood inflow, or in studies using treatments for etiology, such as alcohol abstinence and antivirals (6, 7, 45). New drugs with different mechanisms of action (i.e., acting by decreasing hepatic resistance) would certainly decrease HVPG, but the association between the change in HVPG and clinical benefit might be different and require validation. This validation could be achieved by incorporating HVPG measurements in phase 3 trials (of drugs other than NSBBs) with clinical endpoints. Thus, if the beneficial effect of a reduction in HVPG was to be validated in a phase 3 trial (i.e. correlated with an improvement in clinical outcome), this would strongly favor the approval of new therapies for PH purely based on their ability to reduce portal pressure, as is the case for therapies for arterial hypertension.

d) Feasibility of RCTs with clinical endpoints in patients with compensated cirrhosis.

The relevance of patient selection

The target population to evaluate drugs that would prevent decompensation is that of patients with cirrhosis and CSPH. However, among these patients the risk of decompensation varies widely.

Indeed, data from the cohort included in the timolol trial showed that for every mmHg increase in baseline HVPG there was a 13% increase in the risk of decompensation (figure 3) (1). This indicates that setting the threshold HVPG for inclusion in the study at a level above that defining CSPH (10 mmHg), for example at 14 mmHg, would result in a higher rate of events in a shorter time period and a lower required sample size. Selecting patients based on HVPG would be feasible, as shown by recent studies (43), but may prove difficult in large phase 3 trials. Examples of potential sample sizes according to the minimum expected effect size, length of follow-up and predicted rates of decompensation in the control group are provided in table 4.

There are several non-invasive tools that could substitute for HVPG in its prediction of clinical outcomes in patients with compensated cirrhosis, including laboratory values and imaging-based techniques such as transient elastography. Unfortunately, these have not yet been well validated and will likely need to be disease specific since the association between these non-invasive tests and

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the risk of decompensation might be quantitatively different among patients with viral, alcoholic and non-alcoholic fatty liver disease.

VI. References


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Table 1. Selected new candidate drugs with a proven effect in reducing portal pressure in pre-clinical studies and in some clinical studies.

<table>
<thead>
<tr>
<th>Drugs</th>
<th>Pre-clinical studies</th>
<th>Purported site of action (intrahepatic, extrahepatic or both)</th>
<th>Effect other than ↓ in PP</th>
<th>Potential deleterious signal</th>
<th>Proof of concept studies</th>
<th>Positive effect other than on HVPG</th>
<th>AEs observed</th>
<th>Phase 3 trials</th>
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<tr>
<td>Statins</td>
<td></td>
<td>Intrahepatic (10, 24, 46)</td>
<td>Antifibrotic (47)</td>
<td>Rhabdomyolysis (24)</td>
<td>Reduced HVPG</td>
<td>Yes (16, 48, 49)</td>
<td>Increased in Indocyanine green clearance (16)</td>
<td>Rhabdomyolysis, liver toxicity (17)</td>
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<td>PDE5-inhibitors</td>
<td></td>
<td>Intrahepatic (50-52)</td>
<td>Antifibrotic (52)</td>
<td>-</td>
<td>+/-(discordant) (53-57)</td>
<td>erectile dysfunction</td>
<td>Hypotension</td>
<td>No</td>
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<td>Sorafenib (anti-VEGF)</td>
<td>Both (58-60)</td>
<td>Both (58-60)</td>
<td>Antifibrotic, antiangiogenic (59)</td>
<td>Hepatic injury (61)</td>
<td>+/- (discordant) (62-64)</td>
<td>Improved MAP</td>
<td>Multiple</td>
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<td>Poorly absorbed antibiotics (Rifaximin, Norfloxacin)</td>
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<td>Decreased bacterial translocation (65)</td>
<td>-</td>
<td>+/- (discordant) (66-70)</td>
<td>Less HE, less decompensation (66, 67)</td>
<td>-</td>
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<td>No</td>
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<td>Caspase inhibitors</td>
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<td>Decrease AST (74)</td>
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<td>FXR-agonists</td>
<td>Both (75-77)</td>
<td>Antifibrotic, decreased bacterial translocation (78-81)</td>
<td>-</td>
<td>Yes (82)</td>
<td>-</td>
<td>No</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>Serelaxin</td>
<td>Intrahepatic (83)</td>
<td>Improved Renal perfusion (84)</td>
<td>-</td>
<td>No data (84)</td>
<td>Improved Renal perfusion (84)</td>
<td>-</td>
<td>No</td>
<td>No</td>
</tr>
<tr>
<td>Enoxaparin</td>
<td>Both (85, 86)</td>
<td>Antifibrotic (85)</td>
<td>-</td>
<td>Ongoing trial (NCT02643212)</td>
<td>Decreased decompensation, less portal vein</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Drug</td>
<td>Liver Location</td>
<td>Additional Effects</td>
<td>-/+ (discordant)</td>
<td>Comments</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>-------------------------------------------</td>
<td>----------------</td>
<td>-------------------</td>
<td>-----------------</td>
<td>--------------------------------------------------------------------------</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Endothelin-A receptor antagonists</td>
<td>Intrahepatic</td>
<td>Antifibrotic (89)</td>
<td>+/- (discordant) (91, 92)</td>
<td>-decreased pulmonary arterial pressure (treatment of portopulmonary hypertension) (91) -decreased mean arterial pressure (91) No</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Thalidomide</td>
<td>Intrahepatic</td>
<td>Antifibrotic, antiangiogenic, reduced systemic inflammation (93, 94)</td>
<td>Yes (95)</td>
<td>Reduced systemic inflammation (95) - -</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Angiotensin II Type 1 receptor blocker</td>
<td>Intrahepatic (96-98)</td>
<td>Antifibrotic (98), Improved renal perfusion as low dose (99)</td>
<td>+/- (discordant) (100-106)</td>
<td>Antifibrotic (100) Arterial hypotension, Renal dysfunction (105, 106) No</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Taurine</td>
<td>Intrahepatic and extrahepatic (107)</td>
<td>Antifibrotic (107)</td>
<td>Yes (108)</td>
<td>- -</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Additional references (from #46 on) are provided as supplementary data

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Table 2: Mechanisms behind altered pharmacokinetics in patients with cirrhosis and portal hypertension

<table>
<thead>
<tr>
<th>Mechanism</th>
<th>Comments</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 Reduced Intrinsic Capacity</td>
<td>Liver disease etiology specific variability in different CYPs may exist - for example, alcohol and infection can affect some CYPs selectively</td>
</tr>
<tr>
<td>- Decreased enzyme levels</td>
<td></td>
</tr>
<tr>
<td>- Decreased activity</td>
<td></td>
</tr>
<tr>
<td>2 Reduction in blood flow</td>
<td>May affect drug disposition through a number of mechanisms.</td>
</tr>
<tr>
<td>3 Shunts</td>
<td>Intrahepatic shunts - may account for up to 65% of hepatic blood flow Spontaneous portosystemic shunts and TIPS can significantly modify</td>
</tr>
<tr>
<td>4 Changes in protein binding</td>
<td>Impacts delivery of drugs to hepatocytes and can also have an impact on pharmacodynamic response</td>
</tr>
<tr>
<td>5 Reduced delivery of oxygen</td>
<td>Oxygen is important for CYP activity</td>
</tr>
<tr>
<td>6 Altered transporter expression and function</td>
<td>Hepatocyte uptake as well as efflux into biliary canaliculi can be affected</td>
</tr>
<tr>
<td>7 Small bowel CYP activity</td>
<td>TIPS significantly reduces small bowel CYP3A activity</td>
</tr>
<tr>
<td>8 Drug-drug interactions</td>
<td>Altered pharmacokinetics of one compound may have secondary effects on the pharmacokinetics of co-administered drugs</td>
</tr>
<tr>
<td>9 Impaired renal function</td>
<td>Pharmacokinetics of renally excreted drugs may be profoundly affected in individuals with cirrhosis</td>
</tr>
<tr>
<td>10 Altered intestinal absorption</td>
<td>There may be diminished absorption of orally administered drugs and may also influence the pharmacokinetics of drugs which are bile excreted due to impaired enterohepatic circulation</td>
</tr>
</tbody>
</table>

**Abbreviations:** CYP: Cytochrome P450; TIPS: Transjugular intrahepatic portosystemic shunts
Table 3. Readouts (positive or negative) to be evaluated in proof-of-concept studies of candidate drugs for portal hypertension.

<table>
<thead>
<tr>
<th>Positive readouts</th>
<th>Negative readouts</th>
</tr>
</thead>
<tbody>
<tr>
<td>➢ Decrease in HVPG</td>
<td>• Decrease in mean arterial pressure / renal perfusion</td>
</tr>
<tr>
<td>• Improvement in liver perfusion/liver synthetic function (breath tests, ICG, Hepquant)</td>
<td>• Decrease in liver perfusion/liver synthetic function (breath tests, ICG)</td>
</tr>
<tr>
<td>• Tendency for improvement in clinical outcomes (including mortality)</td>
<td>• Evidence of hepatotoxicity</td>
</tr>
<tr>
<td>• Results support a mechanism of action</td>
<td>• Drug interactions</td>
</tr>
<tr>
<td>• Definition of determinants of response</td>
<td>• Other adverse events (low tolerance)</td>
</tr>
<tr>
<td>• Improvement in patient reported outcomes</td>
<td>• Applicability</td>
</tr>
</tbody>
</table>
**Table 4: Examples of approximate sample sizes** that would be needed in trials for the prevention of decompensation, according to the minimum effect size (which determines the number of events needed), and the rate of decompensation and length of follow-up (which determines the total sample size needed to achieve that number of events). Calculations were performed as proposed by Schoenfeld (Biometrics 1983;39:499-503)

<table>
<thead>
<tr>
<th>Minimum effect size (HR)</th>
<th>N of events needed</th>
<th>Median follow-up (yrs)</th>
<th>Rate decompensation per year in the control group</th>
<th>Total sample size</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.7</td>
<td>247</td>
<td>1</td>
<td>15%</td>
<td>~1500</td>
</tr>
<tr>
<td>0.7</td>
<td>247</td>
<td>2</td>
<td>15%</td>
<td>1000</td>
</tr>
<tr>
<td>0.7</td>
<td>247</td>
<td>2</td>
<td>20%</td>
<td>~700</td>
</tr>
<tr>
<td>0.6</td>
<td>120</td>
<td>1</td>
<td>15%</td>
<td>~1000</td>
</tr>
<tr>
<td>0.6</td>
<td>120</td>
<td>2</td>
<td>15%</td>
<td>~500</td>
</tr>
<tr>
<td>0.6</td>
<td>120</td>
<td>2</td>
<td>20%</td>
<td>~380</td>
</tr>
</tbody>
</table>
Figure legends:

**Fig 1**: Different approaches to treat portal hypertension in cirrhosis. Strategies to decrease portal pressure include drugs that act on intrahepatic targets: fibrosis, endothelial dysfunction, inflammation and drugs that act on decreasing portal flow: splanchnic vasculature, microbiome, and others such as the nervous system, fat, etc as recently reviewed (109,110). A single drug may be able to affect more than one target (multitargeted approach): either two or more intrahepatic targets (e.g. statins) or one intrahepatic and one extrahepatic target (e.g. carvedilol). If a multitargeted drugs is associated with unfavorable effects, the selective delivery either to the intrahepatic or to the extrahepatic circulation may be preferable. Drug therapy should also be tailored to the stage of cirrhosis. In compensated cirrhosis, the main treatment should target the underlying cause of cirrhosis in order to prevent progression of disease (or even to induce regression). Additionally, antifibrotic strategies together with drugs to decrease portal pressure (if significant portal hypertension is present) are beneficial in compensated cirrhosis, since fibrosis and portal hypertension are also self-maintaining processes. In decompensated cirrhosis (ascites, variceal hemorrhage), the main and short-mid-term goal should be reduction in portal pressure and prevention of further complications, since antifibrotic and etiological therapies, although also important, might not have the necessary short-term impact. HSC=hepatic stellate cells; IC= inflammatory cells; KC=Kupffer cells; SEC=sinusoidal endothelial cells.

**Fig 2**: Example of a potential exploratory ordinal outcome including the development of varices. The development of varices would be hierarchically the less severe endpoint, followed by a first decompensation with only one event, and a first decompensation with more than one simultaneous decompensating event (i.e. variceal hemorrhage plus ascites or plus encephalopathy), which is associated with a much higher risk of death than a single decompensating event (41). Death (without decompensation) would be the most severe outcome. The analysis would test whether the study drug induces a significant “category shift” as compared to the control arm.

**Fig 3**: Nomogram showing the 1,2 and 3 yr decompensation risk according to baseline HVPG (constructed with data published in Ripoll et al (1); details are provided in Supplementary File 1). Note the very different incidence of decompensation over time with increasing baseline HVPG values.