

Developed by









Long-Acting Injectable Solid Drug Nanoparticle (SDN) Platform

Verified by the innovator, on Apr 2022

Supported by

Developer(s)

Tandem Nano Ltd. https://www.tandemnano.com/



United Kingdom

A University of Liverpool based start-up company using proprietary technology for the generation of novel nanoparticle formulations to improve the delivery of poorly water soluble APIs.





Unitaid https://unitaid.org

Partnerships

No partner indicated

Technology information

Type of technology

Aqueous drug particle suspension

Administration route

To be determined

Development state and regulatory approval

Active Pharmaceutical Ingredient (API)

Anti-infectives for systemic use, Glecaprevir and pibrentasvir (G/P), Rifapentine

Development Stage

Pre-clinical

Regulatory Approval

Description

The use of particle processing technology to generate long-acting injectable nanoparticle formulations for long-acting delivery.

Technology highlight

The generation of high drug-loading nanoparticles with prolonged release for poorly water-soluble drugs with the potential for co-formulation strategies.

Technology main components

Active pharmaceutical ingredients, FDA/CDER listed excipients.

Delivery device(s)

No delivery device

APIs compatibility profile

API desired features

Water-insoluble molecules

Unit: mg/mL To be determined.

Small molecules

Multiple.

Additional solubility data

None.

_

Additional stability data

API loading: Maximum drug quantity to be loaded

75-90 wt%

API co-administration

2 different APIs : API dependent.

LogP

Scale-up and manufacturing prospects

Scale-up prospects

To be determined

Tentative equipment list for manufacturing

To be determined

Manufacturing

To be determined

Specific analytical instrument required for characterization of formulation

To be determined

Clinical trials

Excipients

Proprietary excipients used

No proprietary excipient used

Novel excipients or existing excipients at a concentration above Inactive Ingredients Database (IID) for the specified route of administration

No novel excipient or existing excipient used

Residual solvents used

No residual solvent used

Additional features

Other features of the technology

Not provided

Release properties

To be determined

Injectability

To be determined

Safety

To be determined

Stability

To be determined

Storage conditions and cold-chain related features

To be determined

Potential application(s)

Therapeutic area(s)

Disease agnostic

Use case(s)

Not provided

Use of technology

Ease of administration

• To be determined

Frequency of administration

Not provided

User acceptance

Targeted user groups

Age Cohort

• Adults

Genders

- All
- Male
- Female
- Cisgender female
- Cisgender male
- Transgender female
- Transgender male
- Intersex
- Gender non-binary

Pregnant individuals

Unspecified

Lactating individuals

Unspecified

Healthy individuals

Unspecified

Comment

Potential associated API(s)

Anti-infectives for systemic use, Glecaprevir and pibrentasvir (G/P), Rifapentine

Class(es)

Not provided

Development stage

Pre-clinical

Clinical trial number(s)

Not provided

Foreseen/approved indication(s)

Not provided

Foreseen user group

Not provided

Foreseen duration between application(s)

Not provided

Applications to Stringent Regulatory Authorities (SRA) / regulatory approvals

Antiparasitic products, Atovaquone

Class(es)

Not provided

Development stage

Pre-clinical

Clinical trial number(s)

Not provided

Foreseen/approved indication(s)

Not provided

Foreseen user group

Not provided

Foreseen duration between application(s)

Not provided

Applications to Stringent Regulatory Authorities (SRA) / regulatory approvals

Patent info

Technology patent families

Patent informations

				Licence	
	Representative			with	Patent
Patent description	patent	Categories	Patent holder	MPP	source
Solid composition comprising	WO2017216564	Compositior	n The Johns Hopkins	Yes	MPP
dispersed atovaquone nanoparticles			University, The		Licence
Expiry date: 2037-06-15			University of Liverpool		
A solid composition comprising					
nanoparticles of atovaquone					
dispersed within one or more					
carrier materials, wherein the					
atovaquone is present in an amount					
of at least 10 wt%. Also described is					
an intramuscularly- or					
subcutaneously-injectable					
formulation of nanoparticles of					
atovaquone					

Patent status/countries	Low, Low- middle and upper-middle	High income
Granted	China, India, Sierra Leone, Eswatini, Liberia, Namibia, Sao Tome and Principe, Mozambique, Zambia, Zimbabwe, Tanzania, United Republic of, Malawi, Ghana, Rwanda, Sudan, Botswana, Lesotho, Kenya, Gambia (the)	Australia, Canada
Filed	Albania, Serbia, Türkiye, North Macedonia, South Africa, Brazil	Liechtenstein, Italy, Norway, Malta, Denmark, Belgium, United Kingdom, Greece, Netherlands, Hungary, Croatia, Switzerland, Spain, San Marino, Slovenia, Austria, Romania, Iceland, Cyprus, Finland, France, Bulgaria, Slovakia, Poland, Latvia, Ireland, Estonia, Germany, Luxembourg, Portugal, Czechia, Lithuania, Monaco, Sweden, Japan, United States of America

Not in force

World Intellectual Property Organization (WIPO), Morocco, Bosnia and Herzegovina, Montenegro, Moldova, Republic of, Uganda World Intellectual Property Organization (WIPO), Chile, United States of America

MPP Licence(s)

Patent and know-how licence on long-acting formulations using Tandem Nano's emulsion-templated freeze-drying technology (ETFD)

https://medicinespatentpool.org/licence-post/long-acting-technologies-for-hcv-tb-and-malaria-treatment

	Representative			Licence with	Patent
Patent description	patent	Categories	Patent holder	MPP	source
Carrier liquids and methods of	WO2013030535	Process	IOTA NANOSOLUTIONS	Yes	MPP
producing such liquids			LIMITED		Licence
Expiry date: 2032-08-20					
The invention provides a method for					
the preparation of a carrier liquid					
which comprises the steps of: (I)					
preparing a single phase solution					
comprising: (a) a solvent or a					
mixture of miscible solvents, (b) a					
liquid carrier material, which is					
soluble in solvent (a), and (c) a					
dopant material which is also					
soluble in solvent (a); (II) cooling					
(preferably freezing) the single					
phase solution produced in step (I)					
to a temperature at which at least					
both the solvent (a) and carrier					
material (b) become solid; and (III)					
removing solid solvent (a) from the					
cooled (frozen) single phase					
solution in vapour form, such that					
the remaining cooled (frozen)					
carrier material (b) and dopant					
material (c) are returned to ambient					
temperature thus providing a					
product of liquid carrier material (b)					
having dopant material (c)					
dispersed therein.					

Patent status/countries	Low, Low- middle and upper-middle	High income
Granted		United Kingdom, Hungary, France,
		Ireland, Germany, United States of
		America
Filed		

Patent status/countries	Low, Low- middle and upper-middle	High income
Not in force	World Intellectual Property Organization (WIPO), Albania, Serbia, Bosnia and Herzegovina, Montenegro, Türkiye, North Macedonia, India	World Intellectual Property Organization (WIPO), Liechtenstein, Italy, Norway, Malta, Denmark, Belgium, United Kingdom, Greece, Netherlands, Croatia, Switzerland, Spain, San Marino, Slovenia, Austria, Romania, Iceland, Cyprus, Finland, Bulgaria, Slovakia, Poland, Latvia, Estonia, Luxembourg, Portugal, Czechia, Lithuania, Monaco,

Sweden

MPP Licence(s)

Patent and know-how licence on long-acting formulations using Tandem Nano's emulsion-templated freeze-drying technology (ETFD)

https://medicinespatentpool.org/licence-post/long-acting-technologies-for-hcv-tb-and-malaria-treatment

				Licence	
	Representative			with	Patent
Patent description	patent	Categories	Patent holder	MPP	source
Nanodispersions of anti-viral drugs	WO2011128623	Composition	n.Duncalf. David John.	Yes	MPP
Expiry date: 2031-04-08		Process	Foster Alison Javne		Licence
The invention provides a			lota Nanosolutions		2.00.000
composition and an antiviral drug			Limited, Long, James,		
preparation, each comprising at			Rannard. Steven Paul.		
least one water-insoluble antiviral			Wang, Dong		
drug and at least one water-soluble					
carrier material, wherein the water-					
insoluble antiviral drug is dispersed					
through the water-soluble carrier					
material in nano-disperse form. The					
present invention further provides					
processes for preparing the					
compositions and drug					
preparations, and also aqueous					
nano-dispersions obtained by					
combining water and the					
compositions.					

Patent status/countries	Low, Low- middle and upper-middle	High income
Granted	India	Liechtenstein, Belgium, United Kingdom, Switzerland, Cyprus, France, Ireland, Germany, Luxembourg, Monaco, Israel, United States of America
Filed		
Not in force	World Intellectual Property Organization (WIPO), China, Albania, Serbia, Bosnia and Herzegovina, Montenegro, Türkiye, North Macedonia	World Intellectual Property Organization (WIPO), Canada, Italy, Norway, Malta, Denmark, United Kingdom, Greece, Netherlands, Hungary, Croatia, Spain, San Marino, Slovenia, Austria, Romania, Iceland, Finland, Bulgaria, Slovakia, Poland, Latvia, Estonia, Portugal, Czechia, Lithuania, Sweden, Japan

MPP Licence(s)

Patent and know-how licence on long-acting formulations using Tandem Nano's emulsion-templated freeze-drying technology (ETFD)

https://medicinespatentpool.org/licence-post/long-acting-technologies-for-hcv-tb-and-malaria-treatment

				Licence	
	Representative			with	Patent
Patent description	patent	Categories	Patent holder	МРР	source
Anti-parasitic nano-dispersed	WO2008006713	Compositior	n Duncalf, David, John,	Yes	MPP
compositions			Essa, Asha, Hassan,		Licence
Expiry date: 2027-06-29			Foster, Alison, Jayne,		
The present invention relates to			Long, James, Rannard,		
nanodisperse antiparasitcs and			Steven, Paul, Unilever		
provides a composition comprising			N.V, Unilever Plc,		
at least one water insoluble anti-			Wang, Dong		
parasitic drug and a water-soluble					
carrier material, wherein the water-					
insoluble anti-parasitic drug					
(preferably an Artemisinin-type					
drug or a quinine type drug) is					
dispersed through the carrier					
material in nano-disperse form					
having a peak diameter of the					

Patent status

nano-disperse form below 1000nm

Patent status/countries	Low, Low- middle and upper-middle	High income
Granted	South Africa, Congo, Mauritania, Guinea-	Canada, Liechtenstein, Italy, Belgium,
	Bissau, Niger, Senegal, Cameroon, Mali,	United Kingdom, Netherlands, Hungary,
	Togo, Burkina Faso, Benin, Côte d'Ivoire,	Croatia, Switzerland, Spain, Austria,
	Central African Republic, Guinea,	France, Ireland, Germany, Sweden,
	Gabon, Equatorial Guinea, Chad	United States of America

Filed

Patent status/countries

Not in force

Low, Low- middle and upper-middle High income

World Intellectual Property Organization (WIPO), Argentina, Brazil, China, Albania, Serbia, Bosnia and Herzegovina, Türkiye, North Macedonia, Mexico, South Africa, India, Sierra Leone, Eswatini, Namibia, Mozambique, Uganda, Zambia, Zimbabwe, Tanzania, United Republic of, Malawi, Ghana, Sudan, Botswana, Lesotho, Kenya, Gambia (the), Indonesia World Intellectual Property Organization (WIPO), Australia, Canada, Chile, Liechtenstein, Italy, Malta, Denmark, Belgium, United Kingdom, Greece, Netherlands, Hungary, Croatia, Switzerland, Spain, Slovenia, Austria, Romania, Iceland, Cyprus, Finland, France, Bulgaria, Slovakia, Poland, Latvia, Ireland, Estonia, Germany, Luxembourg, Portugal, Czechia, Lithuania, Monaco, Sweden, Japan, United States of America, Israel

MPP Licence(s)

Patent and know-how licence on long-acting formulations using Tandem Nano's emulsion-templated freeze-drying technology (ETFD)

https://medicinespatentpool.org/licence-post/long-acting-technologies-for-hcv-tb-and-malaria-treatment

Glecaprevir/Pibrentasvir (LAI candidate)

Patent informations

				Licence	
	Representative			with	Patent
Patent description	patent	Categories Patent	holder	MPP	source
Pibrentasvir compound II	WO2012116257	Compound Abbvie	Inc	No	UNITAID
Expiry date: 2032-02-24					2017
Compounds effective in inhibiting					patent
replication of Hepatitis C virus					landscape
("HCV") are described. This					
invention also relates to processes					
of making such compounds,					
compositions comprising such					
compounds, and methods of using					
such compounds to treat HCV					
infection.					

Patent status/countries	Low, Low- middle and upper-middle	High income
Granted	China, Mexico	Taiwan, Province of China, Spain, Germany, France, United Kingdom, Italy
Filed		Spain
Not in force	World Intellectual Property Organization (WIPO), Türkiye, North Macedonia, Albania, Bosnia and Herzegovina, Montenegro, Serbia	Canada, Japan, United States of America, World Intellectual Property Organization (WIPO), Belgium, Luxembourg, Netherlands, Switzerland, Sweden, Austria, Liechtenstein, Greece, Denmark, Monaco, Portugal, Ireland, Finland, Cyprus, Bulgaria, Czechia, Estonia, Slovakia, Hungary, Poland, Iceland, Malta, Norway, San Marino, Croatia, Romania, Latvia, Lithuania, Slovenia

				Licence	
	Representative			with	Patent
Patent description	patent	Categories	Patent holder	MPP	source
Glecaprevir compound	WO2012040167	Compound	Enanta	Yes	UNITAID
Expiry date: 2031-09-20			Pharmaceuticals, Inc		2017
The present invention discloses					patent
compounds of Formula (I) or					landscape,
pharmaceutically acceptable salts,					MPP
esters, or prodrugs thereof: Formula					Licence,
(I) which inhibit serine protease					Health
activity, particularly the activity of					Canada,
hepatitis C virus (HCV) NS3-NS4A					US FDA
protease. Consequently, the					
compounds of the present invention					
interfere with the life cycle of the					
hepatitis C virus and are also useful					
as antiviral agents. The present					
invention further relates to					
pharmaceutical compositions					
comprising the aforementioned					
compounds for administration to a					
subject suffering from HCV					
infection. The invention also relates					
to methods of treating an HCV					
infection in a subject by					
administering a pharmaceutical					
composition comprising the					
compounds of the present					
invention.					

Patent status

Patent status/countries

Low, Low- middle and upper-middle High income

Granted	Argentina, Brazil, China, Colombia, Dominican Republic, Turkmenistan, Belarus, Tajikistan, Kazakhstan, Azerbaijan, Kyrgyzstan, Armenia, Moldova, Republic of, Ecuador, Türkiye, North Macedonia, Albania, Bosnia and Herzegovina, Montenegro, Serbia, Guatemala, Mexico, Peru, South Africa, India, Bolivia (Plurinational State of), Mongolia, Philippines, Malaysia, Pakistan, Indonesia, Ukraine	Canada, Australia, Cyprus, Denmark, Spain, Hong Kong, Croatia, Israel, Japan, Korea, Republic of, New Zealand, Portugal, Singapore, Slovenia, San Marino, United States of America, Chile, Costa Rica, Russian Federation, Belgium, Germany, France, Luxembourg, Netherlands, Switzerland, United Kingdom, Sweden, Italy, Austria, Liechtenstein, Greece, Monaco, Ireland, Finland, Bulgaria, Czechia, Estonia, Slovakia, Hungary, Poland, Iceland, Malta, Norway, Romania, Latvia, Lithuania, Uruguay, Panama, Bahrain, Kuwait, Qatar, Saudi Arabia, Oman, United Arab Emirates, Macao
Filed	Argentina, Paraguay, Viet Nam, Venezuela (Bolivarian Republic of), Thailand	Cyprus, Denmark, Spain, Croatia, Portugal, Slovenia, San Marino, Taiwan, Province of China, Luxembourg, Netherlands, Hungary, Poland, Norway, Lithuania, Bahrain, Kuwait, Qatar, Saudi Arabia, Oman, United Arab Emirates
Not in force	World Intellectual Property Organization (WIPO), Colombia, Dominican Republic, Ecuador, Türkiye, North Macedonia, Albania, Bosnia and Herzegovina, Montenegro, Serbia, Guatemala, India, Egypt, Malaysia, Indonesia	Australia, Cyprus, Denmark, Spain, Croatia, Japan, Korea, Republic of, Portugal, Slovenia, San Marino, United States of America, World Intellectual Property Organization (WIPO), Costa Rica, Belgium, Germany, France, Luxembourg, Netherlands, Switzerland, United Kingdom, Sweden, Italy, Austria, Liechtenstein, Greece, Monaco, Ireland, Finland, Bulgaria, Czechia, Estonia, Slovakia, Hungary, Poland, Iceland, Malta, Norway, Romania, Latvia, Lithuania, Uruguay, Bahrain, Kuwait, Qatar, Saudi Arabia, Oman, United Arab Emirates

MPP Licence(s)

MPP licence on Glecaprevir/Pibrentasvir (G/P)

https://medicinespatentpool.org/licence-post/glecaprevir-pibrentasvir-g-p/

Patent description	Representative patent	Categories	Patent holder	Licence with MPP	Patent source
Pibrentasvir use in HCV Expiry date: 2033-09-17 Pan-genotypic HCV inhibitors are described. This invention also relates to methods of using these inhibitors to treat HCV infection.	WO2014047039	Use	Abbvie Inc	Yes	MPP Licence

Patent status/countries	Low, Low- middle and upper-middle	High income
Granted	Brazil, Mexico, South Africa, Türkiye, North Macedonia, Albania, Bosnia and Herzegovina, Montenegro, Serbia	Australia, Japan, New Zealand, Belgium, Germany, France, Luxembourg, Netherlands, Switzerland, United Kingdom, Sweden, Italy, Austria, Liechtenstein, Greece, Spain, Denmark, Monaco, Portugal, Ireland, Finland, Cyprus, Bulgaria, Czechia, Estonia, Slovakia, Hungary, Poland, Iceland, Malta, Norway, San Marino, Croatia, Romania, Latvia, Lithuania, Slovenia
Filed	Türkiye, North Macedonia, Albania, Serbia	Canada, Hong Kong, Singapore, Taiwan, Province of China, Belgium, Germany, France, Luxembourg, Netherlands, Switzerland, United Kingdom, Sweden, Italy, Austria, Liechtenstein, Greece, Spain, Denmark, Monaco, Portugal, Ireland, Finland, Cyprus, Bulgaria, Czechia, Estonia, Slovakia, Hungary, Poland, Iceland, Malta, Norway, San Marino, Croatia, Romania, Latvia, Lithuania, Slovenia
Not in force	World Intellectual Property Organization (WIPO), China, Mexico, Bosnia and Herzegovina, Montenegro	Japan, United States of America, World Intellectual Property Organization (WIPO), Russian Federation

https://medicinespatentpool.org/licence-post/glecaprevir-pibrentasvir-g-p/

				Licence	
	Representative			with	Patent
Patent description	patent	Categories	Patent holder	MPP	source
Glecaprevir/Pibrentasvir use in HCV	WO2014152514	llso	Abbyie Inc	Yes	MPP
(without IEN or DDV)	002014152514	036	Abbvie inc	165	Liconco
					LICENCE
Expiry date: 2034-03-14					
The present invention features					
interferon- and ribavirin-free					
therapies for the treatment of HCV.					
Preferably, the treatment is over a					
shorter duration of treatment, such					
as no more than 12 weeks. In one					
aspect, the treatment comprises					
administering at least two direct					
acting antiviral agents without					
interferon and ribavirin to a subject					
with HCV infection, wherein the					
treatment lasts for 12 weeks, and					
said at least two direct acting					
antiviral agents comprise (a)					
Compound 1 or a pharmaceutically					
acceptable salt thereof and (b)					
Compound 2 or a pharmaceutically					
acceptable salt thereof.					

Patent status/countries	Low, Low- middle and upper-middle	High income
Granted	Brazil, Mexico, Serbia, South Africa,	Canada, Australia, Cyprus, Denmark,
	Turkmenistan, Belarus, Tajikistan,	Spain, Israel, Japan, Korea, Republic of,
	Kazakhstan, Azerbaijan, Kyrgyzstan,	New Zealand, Poland, Portugal,
	Armenia, Türkiye, North Macedonia,	Slovenia, Belgium, Germany, France,
	Albania	Luxembourg, Netherlands, Switzerland,
		Russian Federation, United Kingdom,
		Sweden, Italy, Austria, Liechtenstein,
		Greece, Monaco, Ireland, Finland,
		Bulgaria, Czechia, Estonia, Slovakia,
		Hungary, Iceland, Malta, Norway, San
		Marino, Croatia, Romania, Latvia,
		Lithuania

Patent status/countries	Low, Low- middle and upper-middle	High income
Filed	Serbia, Türkiye, North Macedonia,	Cyprus, Denmark, Spain, Hong Kong,
	Albania	Korea, Republic of, Poland, Portugal,
		Singapore, Slovenia, Belgium, Germany,
		France, Luxembourg, Netherlands,
		Switzerland, United Kingdom, Sweden,
		Italy, Austria, Liechtenstein, Greece,
		Monaco, Ireland, Finland, Bulgaria,
		Czechia, Estonia, Slovakia, Hungary,
		Iceland, Malta, Norway, San Marino,
		Croatia, Romania, Latvia, Lithuania
Not in force	World Intellectual Property Organization	Cyprus, Denmark, Spain, Japan, Poland,
	(WIPO), China, Mexico, Serbia,	Portugal, Slovenia, Taiwan, Province of
	Turkmenistan, Belarus, Tajikistan,	China, United States of America, World
	Kazakhstan, Azerbaijan, Kyrgyzstan,	Intellectual Property Organization
	Armenia, Türkiye, North Macedonia,	(WIPO), Belgium, Germany, France,
	Albania, Bosnia and Herzegovina,	Luxembourg, Netherlands, Switzerland,
	Montenegro	Russian Federation, United Kingdom,
		Sweden, Italy, Austria, Liechtenstein,
		Greece, Monaco, Ireland, Finland,
		Bulgaria, Czechia, Estonia, Slovakia,
		Hungary, Iceland, Malta, Norway, San
		Marino, Croatia, Romania, Latvia,

Lithuania

MPP Licence(s)

MPP licence on Glecaprevir/Pibrentasvir (G/P)

https://medicinespatentpool.org/licence-post/glecaprevir-pibrentasvir-g-p/

	Representative			Licence with	Patent
Patent description	patent	Categories	Patent holder	MPP	source
Glecaprevir/Pibrentasvir use in HCV (without IFN or RBV) II Expiry date: 2035-04-01 The present invention features interferon-free therapies for the treatment of HCV. Preferably, the treatment is over a shorter duration of treatment, such as no more than 12 weeks. In one aspect, the treatment comprises administering at least two direct acting antiviral agents to a subject with HCV infection, wherein the treatment lasts for 12 weeks and does not include administration of either interferon or ribavirin, and said at least two direct acting antiviral agents comprise (a) Compound 1 or a pharmaceutically acceptable salt thereof and (b) Compound 2 or a pharmaceutically acceptable salt	WO2015153793	Use	Abbvie Inc	No	UNITAID 2017 patent landscape, US FDA

Patent status/countries	Low, Low- middle and upper-middle	High income
Granted	Mexico	Australia, Japan, United States of America
Filed	China, Albania, North Macedonia, Serbia, Türkiye	Canada, Belgium, Germany, France, Luxembourg, Netherlands, Switzerland, United Kingdom, Sweden, Italy, Austria, Liechtenstein, Greece, Finland, Hungary, Iceland, Ireland, Norway, Poland, Portugal, Romania, San Marino, Bulgaria, Croatia, Cyprus, Czechia, Denmark, Estonia, Latvia, Lithuania, Malta, Monaco, Slovakia, Slovenia, Spain

Not in force

World Intellectual Property Organization (WIPO), China, Bosnia and Herzegovina, Montenegro, Brazil Australia, Japan, United States of America, World Intellectual Property Organization (WIPO)

Patent description	Representative patent	Categories	Patent holder	Licence with MPP	Patent source
Patent description Glecaprevir/Pibrentasvir use in HCV (without IFN or RBV) - treatment regimen Expiry date: 2038-02-09 The present invention features interferon-free therapies for the treatment of HCV. Preferably, the treatment is over a shorter duration of treatment, such as no more than 12 weeks. In one aspect, the treatment comprises administering at least two direct acting antiviral agents to a subject with HCV infection, wherein the treatment lasts for 12 weeks and does not include administration of either	Representative patent CA2994496	Categories	Abbvie Inc	with MPP Yes	Patent source
Interferon or ribavirin, and said at least two direct acting antiviral agents comprise (a) Compound 1 or a pharmaceutically acceptable salt thereof and (b) Compound 2 or a pharmaceutically acceptable salt thereof					

Patent status/countries	Low, Low- middle and upper-middle	High income
Granted		United States of America
Filed		Canada

Not in force

Low, Low- middle and upper-middle High income

China, Brazil, Mexico, Türkiye, North Macedonia, Albania, Bosnia and Herzegovina, Montenegro, Serbia, Moldova, Republic of, Morocco, Tunisia Australia, Japan, United States of America, Belgium, Germany, France, Luxembourg, Netherlands, Switzerland, United Kingdom, Sweden, Italy, Austria, Liechtenstein, Greece, Spain, Denmark, Monaco, Portugal, Ireland, Finland, Cyprus, Bulgaria, Czechia, Estonia, Slovakia, Hungary, Poland, Iceland, Malta, Norway, San Marino, Croatia, Romania, Latvia, Lithuania, Slovenia

MPP Licence(s)

MPP licence on Glecaprevir/Pibrentasvir (G/P)

https://medicinespatentpool.org/licence-post/glecaprevir-pibrentasvir-g-p/

	_			Licence	_
	Representative			with	Patent
Patent description	patent	Categories	Patent holder	мрр	source
Glecaprevir/Pibrentasvir and RBV	WO2014152635	Use	Abbvie Inc	Yes	MPP
use in HCV (without IFN)					Licence
Expiry date: 2034-03-14					
The present invention features					
interferon -free therapies for the					
treatment of HCV. Preferably, the					
treatment is over a shorter duration					
of treatment, such as no more than					
12 weeks. In one aspect, the					
treatment comprises administering					
at least two direct acting antiviral					
agents and ribavirin to a subject					
with HCV infection, wherein the					
treatment lasts for 12 weeks and					
does not include administration of					
interferon, and said at least two					
direct acting antiviral agents					
comprise (a) Compound 1 and (b)					
Compound 2 or a pharmaceutically					
acceptable salt thereof as disclosed					
in the description.					

Patent status/countries	Low, Low- middle and upper-middle	High income
Granted	Serbia, South Africa	Israel, Korea, Republic of
Filed		Canada, Denmark, Spain, Hong Kong,
		Croatia, Israel, Poland, Portugal,
		Singapore, Slovenia, Taiwan, Province of
		China, Norway, Cyprus, San Marino

Patent status/countries

Not in force

Low, Low- middle and upper-middle High income

World Intellectual Property Organization (WIPO), Brazil, China, Mexico, Serbia, Turkmenistan, Belarus, Tajikistan, Kazakhstan, Azerbaijan, Kyrgyzstan, Armenia, Türkiye, North Macedonia, Albania, Bosnia and Herzegovina, Montenegro Australia, Denmark, Spain, Hong Kong, Croatia, Japan, New Zealand, Poland, Portugal, Slovenia, Taiwan, Province of China, United States of America, World Intellectual Property Organization (WIPO), Russian Federation, Norway, Cyprus, Belgium, Germany, France, Luxembourg, Netherlands, Switzerland, United Kingdom, Sweden, Italy, Austria, Liechtenstein, Greece, Monaco, Ireland, Finland, Bulgaria, Czechia, Estonia, Slovakia, Hungary, Iceland, Malta, San Marino, Romania, Latvia, Lithuania

MPP Licence(s)

MPP licence on Glecaprevir/Pibrentasvir (G/P)

https://medicinespatentpool.org/licence-post/glecaprevir-pibrentasvir-g-p/

				Licence	
	Representative			with	Patent
Patent description	patent	Categories	Patent holder	МРР	source
Glecaprevir/Pibrentasvir and RBV	WO2015153792	Use	Abbvie Inc	No	UNITAID
use in HCV (without IFN) II					2017
Expiry date: 2035-04-01					patent
The present invention features					landscape
interferon-free therapies for the					
treatment of HCV. Preferably, the					
treatment is over a shorter duration					
of treatment, such as no more than					
12 weeks. In one aspect, the					
treatment comprises administering					
at least two direct acting antiviral					
agents and ribavirin to a subject					
with HCV infection, wherein the					
treatment lasts for 12 weeks and					
does not include administration of					
interferon, and said at least two					
direct acting antiviral agents					
comprise (a) Compound 1 or a					
pharmaceutically acceptable salt					
thereof and (b) Compound 2 or a					
pharmaceutically acceptable salt					
thereof.					

Patent status/countries	Low, Low- middle and upper-middle	High income
Granted		
Filed		Taiwan, Province of China

Not in force

World Intellectual Property Organization (WIPO), China, Mexico, Albania, North Macedonia, Serbia, Türkiye, Bosnia and Herzegovina, Montenegro Australia, Canada, Japan, United States of America, World Intellectual Property Organization (WIPO), Belgium, Germany, France, Finland, Greece, Hungary, Iceland, Ireland, Italy, Netherlands, Norway, Poland, Portugal, Romania, San Marino, Austria, Bulgaria, Croatia, Cyprus, Czechia, Denmark, Estonia, Latvia, Liechtenstein, Lithuania, Luxembourg, Malta, Monaco, Slovakia, Slovenia, Spain, Sweden, Switzerland, United Kingdom

			Licence	
	Representative		with	Patent
Patent description	patent	Categories Patent holder	MPP	source
Glecaprevir/Pibrentasvir solid	WO2016210273	Composition Abbyie Inc	Yes	MPP
compositions I				Licence
Expiry date: 2036-06-24				
The present invention features solid				
pharmaceutical compositions				
comprising Compound 1 and				
Compound 2. In one embodiment,				
the solid pharmaceutical				
composition includes (1) a first				
layer which comprises 100 mg				
Compound 1, as well as a				
pharmaceutically acceptable				
hydrophilic polymer and a				
pharmaceutically acceptable				
surfactant, all of which are				
formulated in amorphous solid				
dispersion; and (2) a second layer				
which comprises 40 mg Compound				
2, as well as a pharmaceutically				
acceptable hydrophilic polymer and				
a pharmaceutically acceptable				
surfactant, all of which are				
formulated in amorphous solid				
dispersion.				

Patent status/countries	Low, Low- middle and upper-middle	High income
Granted	Mexico, South Africa, Mongolia, Malaysia, Colombia	Australia, Israel, Japan, Korea, Republic of, United States of America, Panama,
		New Zealand

Patent status/countries	Low, Low- middle and upper-middle	High income
Filed	Brazil, Türkiye, India, Ecuador, Guatemala, Thailand, Albania, North Macedonia, Serbia, Bosnia and Herzegovina, Montenegro	Canada, Costa Rica, Belgium, Germany, France, Luxembourg, Netherlands, Switzerland, United Kingdom, Sweden, Italy, Austria, Liechtenstein, Greece, Spain, Denmark, Monaco, Portugal, Ireland, Finland, Cyprus, Bulgaria, Czechia, Estonia, Slovakia, Hungary, New Zealand, Singapore, Hong Kong, Iceland, Norway, Poland, Romania, San Marino, Croatia, Latvia, Lithuania, Malta, Slovenia
Not in force	World Intellectual Property Organization (WIPO), Philippines, China, Dominican Republic, Peru, Turkmenistan, Belarus, Tajikistan, Kazakhstan, Azerbaijan, Kyrgyzstan, Armenia, Egypt, Indonesia, Viet Nam, Ukraine	Japan, United States of America, World Intellectual Property Organization (WIPO), Chile, Russian Federation

MPP Licence(s)

MPP licence on Glecaprevir/Pibrentasvir (G/P)

https://medicinespatentpool.org/licence-post/glecaprevir-pibrentasvir-g-p/

			Licence	
	Representative		with	Patent
Patent description	patent	Categories Patent holder	MPP	source
Glecaprevir/Pibrentasvir solid	WO2017015211	Composition Abbyie Inc	Yes	MPP
compositions II			100	Licence
Expiry date: 2036-07-18				
The present invention features solid				
pharmaceutical compositions				
comprising Compound 1 and				
Compound 2. In one embodiment,				
the solid pharmaceutical				
composition includes (1) a first				
layer which comprises 100 mg				
Compound 1, as well as a				
pharmaceutically acceptable				
hydrophilic polymer and a				
pharmaceutically acceptable				
surfactant, all of which are				
formulated in amorphous solid				
dispersion; and (2) a second layer				
which comprises 40 mg Compound				
2, as well as a pharmaceutically				
acceptable hydrophilic polymer and				
a pharmaceutically acceptable				
surfactant, all of which are				
formulated in amorphous solid				
dispersion.				

Patent status/countries	Low, Low- middle and upper-middle	High income
Granted	South Africa	Australia, Canada, Japan, Israel, New Zealand, Panama

Patent status/countries	Low, Low- middle and upper-middle	High income
Filed	Türkiye, North Macedonia, Albania,	Korea, Republic of, Costa Rica, Belgium,
	Bosnia and Herzegovina, Montenegro,	Germany, France, Luxembourg,
	Serbia, Ecuador, Guatemala, Mongolia,	Netherlands, Switzerland, United
	Thailand	Kingdom, Sweden, Italy, Austria,
		Liechtenstein, Greece, Spain, Denmark,
		Monaco, Portugal, Ireland, Finland,
		Cyprus, Bulgaria, Czechia, Estonia,
		Slovakia, Hungary, Poland, Iceland,
		Malta, Norway, San Marino, Croatia,
		Romania, Latvia, Lithuania, Slovenia,
		New Zealand, Singapore, Hong Kong
Not in force	World Intellectual Property Organization	Korea, Republic of, United States of
	(WIPO), Brazil, China, Colombia,	America, World Intellectual Property
	Philippines, Peru, Dominican Republic,	Organization (WIPO), Chile, Russian
	Turkmenistan, Belarus, Tajikistan,	Federation
	Kazakhstan, Egypt, Indonesia, Viet Nam,	
	India, Mexico, Moldova, Republic of,	
	Malaysia, Ukraine	

MPP Licence(s)

MPP licence on Glecaprevir/Pibrentasvir (G/P)

https://medicinespatentpool.org/licence-post/glecaprevir-pibrentasvir-g-p/

Patent description	Representative patent	Categories	Patent holder	Licence with MPP	Patent source
Pibrentasvir compound Expiry date: 2031-10-12 Compounds effective in inhibiting replication of Hepatitis C virus (HCV) are described. This invention also relates to processes of making such compounds, compositions comprising such compounds, and	WO2012051361	Compound	Abbott Laboratories	Yes	Health Canada, US FDA, MPP Licence
methods of using such compounds					

Patent status

to treat HCV infection.

Patent status/countries	Low, Low- middle and upper-middle	High income
Granted	Colombia, Argentina, China, Dominican	United States of America, Australia,
	Republic, Turkmenistan, Belarus,	Chile, Japan, Korea, Republic of, New
	Tajikistan, Kazakhstan, Azerbaijan,	Zealand, Singapore, Taiwan, Province of
	Kyrgyzstan, Armenia, Moldova, Republic	China, Uruguay, Denmark, Spain,
	of, Ecuador, Türkiye, North Macedonia,	Portugal, Slovenia, Canada, Israel, Hong
	Albania, Bosnia and Herzegovina,	Kong, Russian Federation, Belgium,
	Montenegro, Serbia, Mexico, Peru,	Germany, France, Luxembourg,
	Ukraine, Bolivia (Plurinational State of),	Netherlands, Switzerland, United
	Indonesia, Malaysia, Philippines, Viet	Kingdom, Sweden, Italy, Austria,
	Nam, South Africa, Brazil	Liechtenstein, Greece, Monaco, Ireland,
		Finland, Cyprus, Bulgaria, Czechia,
		Estonia, Slovakia, Hungary, Poland,
		Iceland, Malta, Norway, Croatia,
		Romania, Latvia, Lithuania, Panama

Patent status/countries

Filed

Low, Low- middle and upper-middle High income

Ecuador, Türkiye, North Macedonia, Albania, Bosnia and Herzegovina, Montenegro, Serbia, India, Bolivia (Plurinational State of), Mongolia, Pakistan, Paraguay, Thailand, Venezuela (Bolivarian Republic of), Guatemala Costa Rica, Denmark, Spain, Portugal, Slovenia, Belgium, Germany, France, Luxembourg, Netherlands, Switzerland, United Kingdom, Sweden, Italy, Austria, Liechtenstein, Greece, Monaco, Ireland, Finland, Cyprus, Bulgaria, Czechia, Estonia, Slovakia, Hungary, Poland, Iceland, Malta, Norway, San Marino, Croatia, Romania, Latvia, Lithuania, Bahrain, Kuwait, Qatar, Saudi Arabia, Oman, United Arab Emirates

Not in force

World Intellectual Property Organization (WIPO), Argentina, China, Turkmenistan, Belarus, Tajikistan, Kazakhstan, Azerbaijan, Kyrgyzstan, Armenia, Moldova, Republic of, Türkiye, North Macedonia, Albania, Bosnia and Herzegovina, Montenegro, Serbia, Mexico, Peru, Egypt, Viet Nam United States of America, World Intellectual Property Organization (WIPO), Chile, Costa Rica, New Zealand, Uruguay, Denmark, Spain, Portugal, Slovenia, Canada, Russian Federation, Belgium, Germany, France, Luxembourg, Netherlands, Switzerland, United Kingdom, Sweden, Italy, Austria, Liechtenstein, Greece, Monaco, Ireland, Finland, Cyprus, Bulgaria, Czechia, Estonia, Slovakia, Hungary, Poland, Iceland, Malta, Norway, San Marino, Croatia, Romania, Latvia, Lithuania

MPP Licence(s)

MPP licence on Glecaprevir/Pibrentasvir (G/P)

https://medicinespatentpool.org/licence-post/glecaprevir-pibrentasvir-g-p/

Patent description	Representative patent	Categories Patent holder	Licence with MPP	Patent source
Glecaprevir crystal forms Expiry date: 2035-06-05 The present invention features crystalline forms of Compound I. In one embodiment, a crystalline form of Compound I has characteristic peaks in the PXRD pattern as shown in any one of Figures 1-4.	WO2015188045	Polymorphs Abbvie Inc	No	US FDA

Patent status/countries	Low, Low- middle and upper-middle	High income
Granted	Mexico	United States of America, Australia
Filed	Türkiye, North Macedonia, Albania, Serbia	Canada, Japan, Belgium, Germany, France, Luxembourg, Netherlands, Switzerland, United Kingdom, Sweden, Italy, Austria, Liechtenstein, Greece, Spain, Denmark, Monaco, Portugal, Ireland, Finland, Cyprus, Bulgaria, Czechia, Estonia, Slovakia, Hungary, Poland, Iceland, Malta, Norway, San Marino, Croatia, Romania, Latvia, Lithuania, Slovenia
Not in force	World Intellectual Property Organization (WIPO), Türkiye, North Macedonia, Albania, Bosnia and Herzegovina, Montenegro, Serbia, Morocco, China	Australia, Japan, World Intellectual Property Organization (WIPO), Belgium, Germany, France, Luxembourg, Netherlands, Switzerland, United Kingdom, Sweden, Italy, Austria, Liechtenstein, Greece, Spain, Denmark, Monaco, Portugal, Ireland, Finland, Cyprus, Bulgaria, Czechia, Estonia, Slovakia, Hungary, Poland, Iceland, Malta, Norway, San Marino, Croatia, Romania, Latvia, Lithuania, Slovenia

	Representative		Licence with	Patent
Patent description	patent	Categories Patent holder	MPP	source
Pibrentasvir crystal forms	WO2015171993	Polymorphs Abbvie Inc	No	UNITAID
Expiry date: 2035-05-08				2017
The present invention features				patent
crystalline forms of Compound I. In				landscape,
one embodiment, a crystalline form				Pat-
of Compound I has characteristic				Informed
peaks in the PXRD pattern as shown				
in one of Figures 1-10.				

Patent status/countries	Low, Low- middle and upper-middle	High income
Granted	Mexico	Australia, Japan, United States of America
Filed	China, Albania, Serbia, Türkiye, North Macedonia	Canada, Liechtenstein, Italy, Norway, Malta, Denmark, Belgium, United Kingdom, Greece, Netherlands, Hungary, Croatia, Switzerland, Spain, San Marino, Slovenia, Austria, Romania, Iceland, Cyprus, Finland, France, Bulgaria, Slovakia, Poland, Latvia, Ireland, Estonia, Germany, Luxembourg, Portugal, Czechia, Lithuania, Monaco, Sweden, United States of America
Not in force	World Intellectual Property Organization (WIPO), China, Morocco, Albania, Serbia, Bosnia and Herzegovina, Montenegro, Türkiye, North Macedonia, Mexico	World Intellectual Property Organization (WIPO), Australia, Liechtenstein, Italy, Norway, Malta, Denmark, Belgium, United Kingdom, Greece, Netherlands, Hungary, Croatia, Switzerland, Spain, San Marino, Slovenia, Austria, Romania, Iceland, Cyprus, Finland, France, Bulgaria, Slovakia, Poland, Latvia, Ireland, Estonia, Germany, Luxembourg, Portugal, Czechia, Lithuania, Monaco, Sweden, Japan

Supporting material

Publications

Improving maraviroc oral bioavailability by formation of solid drug nanoparticles. Savage AC, Tatham LM, Siccardi M, Scott T, Vourvahis M, Clark A, Rannard SP, Owen A. Eur J Pharm Biopharm. 2019 May;138:30-36. span style="color: rgb(33, 33, 33);">doi: 10.1016/j.ejpb.2018.05.015.

Oral drug administration remains the preferred approach for treatment of HIV in most patients. Maraviroc (MVC) is the first in class co-receptor antagonist, which blocks HIV entry into host cells. MVC has an oral bioavailability of approximately 33%, which is limited by poor permeability as well as affinity for CYP3A and several drug transporters. While once-daily doses are now the favoured option for HIV therapy, dose-limiting postural hypotension has been of theoretical concern when administering doses high enough to achieve this for MVC (particularly during coadministration of enzyme inhibitors). To overcome low bioavailability and modify the pharmacokinetic profile, a series of 70 wt% MVC solid drug nanoparticle (SDN) formulations (containing 30 wt% of various polymer/surfactant excipients) were generated using emulsion templated freeze-drying. The lead formulation contained PVA and AOT excipients (MVCSDNPVA/AOT), and was demonstrated to be fully water-dispersible to release drug nanoparticles with z-average diameter of 728 nm and polydispersity index of 0.3. In vitro and in vivo studies of MVCSDNPVA/AOT showed increased apparent permeability of MVC, compared to a conventional MVC preparation, with in vivo studies in rats showing a 2.5-fold increase in AUC (145.33 vs. 58.71 ng h ml-1). MVC tissue distribution was similar or slightly increased in tissues examined compared to the conventional MVC preparation, with the exception of the liver, spleen and kidneys, which showed statistically significant increases in MVC for MVCSDNPVA/AOT. These data support a novel oral format with the potential for dose reduction while maintaining therapeutic MVC exposure and potentially enabling a once-daily fixed

dose combination product.

Antiretroviral solid drug nanoparticles with enhanced oral bioavailability: production, characterization, and in vitro-in vivo correlation. McDonald TO, Giardiello M, Martin P, Siccardi M, Liptrott NJ, Smith D, Roberts P, Curley P, Schipani A, Khoo SH, Long J, Foster AJ, Rannard SP, Owen A. span style="color: rgb(33, 33, 33);">Adv Healthc Mater. 2014 Mar;3(3):400-11. span style="color: rgb(33, 33, 33);">doi: 10.1002/adhm.201300280.

Nanomedicine strategies have produced many commercial products. However, no orally dosed HIV nanomedicines are available clinically to patients. Although nanosuspensions of drug particles have demonstrated many benefits, experimentally achieving >25 wt% of drug relative to stabilizers is highly challenging. In this study, the emulsion-templated freeze-drying technique for nanoparticles formation is applied for the first time to optimize a nanodispersion of the leading non-nucleoside reverse transcriptase inhibitor efavirenz, using clinically acceptable polymers and surfactants. Dry monoliths containing solid drug nanoparticles with extremely high drug loading (70 wt% relative to polymer and surfactant stabilizers) are stable for several months and reconstitute in aqueous media to provide nanodispersions with *z*-average diameters of 300 nm. The solid drug nanoparticles exhibit reduced cytoxicity and increased in vitro transport through model gut epithelium. In vivo studies confirm bioavailability benefits with an approximately four-fold higher pharmacokinetic exposure after oral administration to rodents, and predictive modeling suggests dose reduction with the new formulation may be possible.

Accelerated oral nanomedicine discovery from miniaturized screening to clinical production exemplified by paediatric HIV nanotherapies. Giardiello M, Liptrott NJ, McDonald TO, Moss D, Siccardi M, Martin P, Smith D, Gurjar R, Rannard SP, Owen A. Nat Commun. 2016 Oct 21;7:13184. ?doi: 10.1038/ncomms13184.

Considerable scope exists to vary the physical and chemical properties of nanoparticles, with subsequent impact on biological interactions; however, no accelerated process to access large nanoparticle material space is currently available, hampering the development of new nanomedicines. In particular, no clinically available nanotherapies exist for HIV populations and conventional paediatric HIV medicines are poorly available; one current paediatric formulation utilizes high ethanol concentrations to solubilize lopinavir, a poorly soluble antiretroviral. Here we apply accelerated nanomedicine discovery to generate a potential aqueous paediatric HIV nanotherapy, with clinical translation and regulatory approval for human evaluation. Our rapid small-scale screening approach yields large libraries of solid drug nanoparticles (160 individual components) targeting oral dose. Screening uses 1 mg of drug compound per library member and iterative pharmacological and chemical evaluation establishes potential candidates for progression through to clinical manufacture. The wide applicability of our strategy has implications for multiple therapy development programmes.

Long-acting injectable atovaquone nanomedicines for malaria prophylaxis. Bakshi, R.P., Tatham, L., Savage, A.C. <em style="color: rgb(34, 34, 34);">et al.et al. <em style="color: rgb(34, 34, 34);">Nat Commun <strong style="color: rgb(34, 34, 34);">9, 315 (2018). <ep>style="color: rgb(34, 34, 34);">315 (2018). ?https://doi.org/10.1038/s41467-017-02603-z

Chemoprophylaxis is currently the best available prevention from malaria, but its

efficacy is compromised by non-adherence to medication. Here we develop a longacting injectable formulation of atovaquone solid drug nanoparticles that confers longlived prophylaxis against *Plasmodium berghei* ANKA malaria in C57BL/6 mice. Protection is obtained at plasma concentrations above 200 ng ml-1 and is causal, attributable to drug activity against liver stage parasites. Parasites that appear after subtherapeutic doses remain atovaquone-sensitive.

Pharmacokinetic-pharmacodynamic analysis indicates protection can translate to humans at clinically achievable and safe drug concentrations, potentially offering protection for at least 1 month after a single administration. These findings support the use of long-acting injectable formulations as a new approach for malaria prophylaxis in travellers and for malaria control in the field.

Additional documents

No documents were uploaded

Useful links

There are no additional links

Access principles

Collaborate for development



Consider on a case by case basis, collaborating on developing long acting products with potential significant public health impact, especially for low- and middle-income countries (LMICs), utilising the referred to long-acting technology

Agree Share technical information for match-making assessment



Provide necessary technical information to a potential partner, under confidentiality agreement, to enable preliminary assessment of whether specific medicines of public health importance in LMICs might be compatible with the referred to long-acting technology to achieve a public health benefit

Agree Work with MPP to expand access in LMICs



In the event that a product using the referred to long-acting technology is successfully developed, the technology IP holder(s) will work with the Medicines Patent Pool towards putting in place the most appropriate strategy for timely and affordable access in low and middle-income countries, including through licensing

Agree

Comment & Information

No.

Illustrations



SDN banner