
UNITED STATES
SECURITIES AND EXCHANGE COMMISSION
Washington, D.C. 20549

FORM 6-K

REPORT OF FOREIGN PRIVATE ISSUER PURSUANT TO RULE 13a-16 OR 15d-16
UNDER THE SECURITIES EXCHANGE ACT OF 1934

For the month of **March 2020**

Commission File Number: **001-38480**

IMV Inc.

(Name of registrant)

130 Eileen Stubbs Avenue, Suite 19 Dartmouth, Nova Scotia B3B 2C4, Canada

Indicate by check mark whether the registrant files or will file annual reports under cover of Form 20-F or Form 40-F.

Form 20-F Form 40-F

Indicate by check mark if the registrant is submitting the Form 6-K in paper as permitted by Regulation S-T Rule 101(b)(1):

Indicate by check mark if the registrant is submitting the Form 6-K in paper as permitted by Regulation S-T Rule 101(b)(7):

SIGNATURES

Pursuant to the requirements of the Securities Exchange Act of 1934, the registrant has duly caused this report to be signed on its behalf by the undersigned, thereunto duly authorized.

IMV Inc.

Date: March 15, 2020

By: /s/ Pierre Labbé
Name: Pierre Labbé
Title: Chief Financial Officer

Form 6-K Exhibit Index

Exhibit Number	Document Description
99.1	Interim Financial Statements for the period ended March 31, 2020
99.2	Management Discussion and Analysis for the period ended March 31, 2020
99.3	CEO certification
99.4	CFO certification



Unaudited Interim Condensed Consolidated Financial Statements

March 31, 2020

May 14, 2020

Management's Responsibility for Financial Reporting

The accompanying unaudited interim condensed consolidated financial statements of **IMV Inc. (the "Corporation")** are the responsibility of management and have been approved by the Board of Directors. The consolidated financial statements have been prepared by management in accordance with International Financial Reporting Standards as issued by the International Accounting Standards Board. The unaudited interim condensed consolidated financial statements include some amounts and assumptions based on management's best estimates which have been derived with careful judgment.

In fulfilling its responsibilities, management has developed and maintains a system of internal accounting controls. These controls are designed to ensure that the financial records are reliable for preparation of the unaudited interim condensed consolidated financial statements. The Audit Committee of the Board of Directors reviewed and approved the Corporation's unaudited interim condensed consolidated financial statements, and recommended their approval by the Board of Directors.

(signed) "*Frederic Ors*"
Chief Executive Officer

(signed) "*Pierre Labb *"
Chief Financial Officer

Approved on behalf of the Board of Directors

(signed) "*James W. Hall*", Director

(signed) "*Wayne Pisano*", Director

IMV Inc.

Unaudited Interim Condensed Consolidated Statements of Financial Position

As at March 31, 2020 and December 31, 2019

(Expressed in thousands of Canadian dollars except for share and per share amounts)

	March 31, 2020	December 31, 2019
	\$	\$
Assets		
Current assets		
Cash and cash equivalents	7,372	14,066
Amounts receivable	653	845
Prepaid expenses	2,676	3,032
Investment tax credits receivable	1,886	1,661
	<u>12,587</u>	<u>19,604</u>
Property and equipment	<u>2,742</u>	<u>2,830</u>
	<u>15,329</u>	<u>22,434</u>
Liabilities		
Current liabilities		
Accounts payable and accrued liabilities	7,525	6,157
Amounts due to directors	64	60
Current portion of long-term debt (note 4)	342	88
Current portion of lease obligation	103	100
	<u>8,034</u>	<u>6,405</u>
Lease obligation	1,181	1,208
Long-term debt (note 4)	<u>8,529</u>	<u>8,373</u>
	<u>17,744</u>	<u>15,986</u>
Equity	<u>(2,415)</u>	<u>6,448</u>
	<u>15,329</u>	<u>22,434</u>

The accompanying notes form an integral part of these unaudited interim condensed consolidated financial statements.

IMV Inc.

Unaudited Interim Condensed Consolidated Statements of Changes in Equity

For the periods ended March 31, 2020 and March 31, 2019

(Expressed in thousands of Canadian dollars except for share and per share amounts)

	Share Capital	Contributed Surplus	Warrants	Deficit	Total
	\$	\$	\$	\$	\$
	(note 5)	(note 6)	(note 7)		
Balance, December 31, 2018	90,152	6,504	415	(92,754)	4,317
Net loss and comprehensive loss for the period	-	-	-	(5,943)	(5,943)
Issuance of shares in public offering	29,456	-	-	-	29,456
Share issuance costs	(2,525)	-	-	-	(2,525)
Exercise of warrants	82	-	(21)	-	61
Employee share options:					
Value of services recognized	-	391	-	-	391
Exercise of options	242	(187)	-	-	55
Balance, March 31, 2019	117,407	6,708	394	(98,697)	25,812
Balance, December 31, 2019	117,544	8,691	332	(120,119)	6,448
Net loss and comprehensive loss for the period	-	-	-	(9,664)	(9,664)
Issuance of shares in "at-the-market" public offering	771	-	-	-	771
Share issuance costs	(426)	-	-	-	(426)
Redemption of deferred share units, net of applicable taxes	184	(189)	-	-	(5)
Expiry of warrants	-	332	(332)	-	-
Deferred share units ("DSU"s):					
Value of services recognized	-	122	-	-	122
Employee share options:					
Value of services recognized	-	306	-	-	306
Exercise of options	239	(206)	-	-	33
Balance, March 31, 2020	118,312	9,056	-	(129,783)	(2,415)

The accompanying notes form an integral part of these unaudited interim condensed consolidated financial statements.

IMV Inc.

Unaudited Interim Condensed Consolidated Statements of Loss and Comprehensive Loss

For the three months ended March 31, 2020 and 2019

(Expressed in thousands of Canadian dollars except for share and per share amounts)

	Three months ended March 31, 2020	Three months ended March 31, 2019
	\$	\$
Income		
Subcontract revenue	-	8
Interest income	68	74
	<u>68</u>	<u>82</u>
Expenses		
Research and development	6,824	4,013
General and administrative	3,033	1,960
Government assistance	(558)	(346)
Accreted interest (note 4)	433	398
	<u>9,732</u>	<u>6,025</u>
Net loss and comprehensive loss for the year	<u>(9,664)</u>	<u>(5,943)</u>
Basic and diluted loss per share	<u>(0.19)</u>	<u>(0.13)</u>
Weighted-average shares outstanding	<u>50,719,488</u>	<u>46,712,436</u>

The accompanying notes form an integral part of these unaudited interim condensed consolidated financial statements.

IMV Inc.

Unaudited Interim Condensed Consolidated Statements of Cash Flows

For the three months ended March 31, 2020 and 2019

(Expressed in thousands of Canadian dollars except for share and per share amounts)

	Three months ended March 31, 2020 \$	Three months ended March 31, 2019 \$
Cash provided by (used in)		
Operating activities		
Net loss and comprehensive loss for the year	(9,664)	(5,943)
Charges to operations not involving cash		
Interest on lease obligation	36	38
Depreciation of property and equipment	121	107
Accretion of long-term debt	433	398
Deferred share unit compensation	122	(203)
Stock-based compensation	306	391
Loss on disposal of assets	-	8
	<u>(8,646)</u>	<u>(5,204)</u>
Net change in non-cash working capital balances related to operations		
Decrease in amounts receivable	192	459
Decrease in prepaid expenses	356	173
Increase in investment tax credits receivable	(225)	(345)
Increase (decrease) in accounts payable and accrued liabilities	1,368	(3,303)
Increase in amounts due to directors	4	14
	<u>(6,951)</u>	<u>(8,206)</u>
Financing activities		
Proceeds from public equity offering	771	29,456
Share issuance costs in public equity offering	(426)	(1,862)
Proceeds from the exercise of stock options	33	55
Proceeds from the exercise of warrants	-	61
Withholdings on redemption of DSUs	(5)	-
Repayment of long-term debt	(23)	(20)
Repayment of lease obligation	(60)	(60)
	<u>290</u>	<u>27,630</u>
Investing activities		
Acquisition of property and equipment	(33)	(112)
Net change in cash and cash equivalents during the period	<u>(6,694)</u>	<u>19,312</u>
Cash and cash equivalents - Beginning of period	<u>14,066</u>	<u>14,895</u>
Cash and cash equivalents - End of period	<u>7,372</u>	<u>34,207</u>
Supplementary cash flow		
Interest received	68	74

The accompanying notes form an integral part of these unaudited interim condensed consolidated financial statements.

IMV Inc.

Notes to the Unaudited Interim Condensed Consolidated Financial Statements

As at March 31, 2020 and December 31, 2019

(Expressed in thousands of Canadian dollars except for share and per share amounts)

1 Nature of operations

IMV Inc. (the “Corporation or IMV”) is, through its 100% owned subsidiary, a clinical stage biopharmaceutical company dedicated to making immunotherapy more effective, more broadly applicable and more widely available to people facing cancer and other serious diseases. IMV is pioneering a new class of immunotherapies based on the Corporation’s proprietary drug delivery platform (“DPX”). This patented technology leverages a novel mechanism of action that enables the programming of immune cells in vivo, which are aimed at generating powerful new synthetic therapeutic capabilities. IMV’s lead candidate, DPX-Survivac, is a T cell-activating immunotherapy that combines the utility of the platform with a target: survivin. IMV is currently assessing DPX-Survivac in advanced ovarian cancer, as well as a combination therapy in multiple clinical studies with Merck’s Keytruda® Checkpoint inhibitor. The Corporation has one reportable and geographic segment. Incorporated under the Canada Business Corporations Act and domiciled in Dartmouth, Nova Scotia, the shares of the Corporation are listed on the Nasdaq Stock Market and the Toronto Stock Exchange under the symbol “IMV”. The address of its principal place of business is 130 Eileen Stubbs Avenue, Suite 19, Dartmouth, Nova Scotia, Canada.

An outbreak of a novel strain of coronavirus, identified as “COVID-19”, was declared a global pandemic by the World Health Organization on March 11, 2020. In response, many countries have required entities to limit or suspend business operations and implemented travel restrictions and quarantine measures. These measures have disrupted the activities of many entities and have led to significant volatility in the global markets. The Corporation continues to monitor and actively manage the developing impacts from COVID-19, including but not limited to, the potential future effects on its assets, cash flow and liquidity, and will continue to assess impacts to the Corporation’s operations, going concern assumption, and the value of assets and liabilities reported in these statements. In addition, a significant outbreak of contagious diseases in the human population could result in a widespread health crisis that could adversely affect the economies and financial markets of many countries, resulting in an economic downturn that could further affect the Company’s operations and ability to finance its operations.

2 Basis of presentation

The Corporation prepares its unaudited interim condensed consolidated financial statements in accordance with Canadian generally accepted accounting principles as set out in the Chartered Professional Accountants of Canada Handbook - Accounting Part I, which incorporates International Financial Reporting Standards (“IFRS”) as issued by the International Accounting Standards Board (“IASB”).

These unaudited interim condensed consolidated financial statements have been prepared in accordance with IFRS applicable to the preparation of interim financial statements, including IAS 34, International Accounting Standards 34 *Interim Financial Reporting*. Accordingly, certain information normally included in annual audited financial statements prepared in accordance with IFRS, as issued by the IASB, have been omitted or condensed. The unaudited interim condensed consolidated financial statements should be read in conjunction with the Corporation’s annual audited consolidated financial statements for the year ended December 31, 2019.

The policies applied in these unaudited interim condensed consolidated financial statements are based on IFRS issued and outstanding as of May 14, 2020, the date the Board of Directors approved the statements. Any subsequent changes to IFRS that are given effect in the Corporation’s annual audited consolidated financial statements for the year ending December 31, 2019 could result in restatement of these unaudited interim condensed consolidated financial statements.

IMV Inc.

Notes to the Unaudited Interim Condensed Consolidated Financial Statements

As at March 31, 2020 and December 31, 2019

(Expressed in thousands of Canadian dollars except for share and per share amounts)

3 Significant accounting policies, judgments and estimation uncertainty

These unaudited interim condensed consolidated financial statements have been prepared using the same policies and methods as the annual audited consolidated financial statements of the Corporation for the year ended December 31, 2019. Refer to note 4 of the Corporation's annual audited consolidated financial statements for the year ended December 31, 2019 for more information on accounting policies and methods applied.

4 Long-term debt

	March 31, 2020	December 31, 2019
	\$	\$
Atlantic Canada Opportunities Agency ("ACOA") Atlantic Innovation Fund interest-free loan with a maximum contribution of \$3,786. Annual repayments, commencing December 1, 2008, are calculated as a percentage of gross revenue for the preceding fiscal year, at 2% when gross revenues are less than \$5,000 and 5% when gross revenues are greater than \$5,000. As at March 31, 2020, the amount drawn down on the loan, net of repayments, is \$3,744 (2019 - \$3,744).	1,527	1,404
ACOA Atlantic Innovation Fund interest-free loan with a maximum contribution of \$3,000. Annual repayments, commencing December 1, 2011, are calculated as a percentage of gross revenue for the preceding fiscal year, at 2% when gross revenues are less than \$5,000 and 5% when gross revenues are greater than \$5,000. As at March 31, 2020, the amount drawn down on the loan is \$2,995 (2019 - \$2,995).	1,345	1,237
ACOA Business Development Program, interest-free loan with a maximum contribution of \$395, repayable in monthly payments beginning October 2015 of \$3 until October 2017 and \$6 until September 2022. As at March 31, 2020, the amount drawn down on the loan, net of repayments, is \$167 (2019 - \$184).	164	180
ACOA Atlantic Innovation Fund interest-free loan with a maximum contribution of \$2,944, annual repayments commencing September 1, 2014, are calculated as a percentage of gross revenue from specific product(s) for the preceding fiscal year, at 5% for the first 5 year period and 10%, thereafter. As at March 31, 2020, the amount drawn down on the loan is \$2,944 (2019 - \$2,944).	1,611	1,481
TNC 120-140 Eileen Stubbs Ltd. (the "Landlord") loan, with a maximum contribution of \$300,000, bearing interest at 8% annum, is repayable in monthly payments beginning upon receipt of the final installment of the loan until May 31, 2028. The loan is made available in three equal installments based on the Corporation meeting certain milestones. As at March 31, 2020, the amount drawn down on the loan is \$273 (2019 - \$279).	273	279
Province of Nova Scotia (the "Province") secured loan with a maximum contribution of \$5,000, bearing interest at a rate equal to the Province's cost of funds plus 1%, compounded semi-annually and payable monthly. The loan is repayable in monthly payments beginning January 1, 2021 of \$83 plus interest until December 2025. The Corporation and its subsidiary have provided a general security agreement granting a first security interest in favour of the Province of Nova Scotia in and to all the assets of the Corporation and its subsidiary, including the intellectual property. As at March 31, 2020, the amount drawn down on the loan is \$5,000 (2019 - \$5,000).	3,951	3,880
	8,871	8,461
Less: Current portion	342	88
	<u>8,529</u>	<u>8,373</u>

(3)

IMV Inc.

Notes to the Unaudited Interim Condensed Consolidated Financial Statements

As at March 31, 2020 and December 31, 2019

(Expressed in thousands of Canadian dollars except for share and per share amounts)

4 Long-term debt (continued)

Total contributions received, less amounts that have been repaid as at March 31, 2020, is \$15,124 (2019 - \$15,147).

Certain ACOA loans and the Province loan require approval by ACOA or the Minister for the Province before the Corporation can pay management fees, bonuses, dividends or other distributions, or before there is any change of ownership of the Corporation. The Province loan requires the Corporation to obtain the written consent of the Province prior to the sale, disposal or abandonment of possession of the intellectual property of the Corporation or its subsidiary. If during the term of the Province loan, the head office, research and development facilities, or production facilities of the Corporation are moved from the Province, the Corporation is required to repay 40% of the outstanding principal of the loan.

In June 2019, the Corporation amended its loan agreement with the Province. Previously, the maturity date of the loan was August 9, 2020. The Corporation shall now start repaying the balance of the principal amount on the first day of January 2021, by making 60 monthly principal payments of \$83 plus interest from January 2021 to December 2025. The annual interest rate remains at the Province's cost of funds plus 1%.

The Province loan requires certain early repayments if the Corporation's subsidiary, or the Corporation on a consolidated basis, has cash flow from operations in excess of \$1,500. The Province loan also requires repayment of the loan under certain circumstances, such as changes of control, sale or liquidation of the Corporation or the sale of substantially all of the assets of the Corporation.

	March 31, 2020	December 31, 2019
	\$	\$
Balance - Beginning of year	8,461	8,150
Accreted interest	433	1,239
Revaluation of long-term debt	-	(840)
Repayment of debt	(23)	(88)
Balance - End of year	8,871	8,461
Less: Current portion	342	88
Non-current portion	8,529	8,373

The Corporation is in compliance with its debt covenants as of March 31, 2020.

IMV Inc.

Notes to the Unaudited Interim Condensed Consolidated Financial Statements

As at March 31, 2020 and December 31, 2019

(Expressed in thousands of Canadian dollars except for share and per share amounts)

5 Share capital**Authorized**

Unlimited number of common shares and preferred shares, issuable in series, all without par value.

	Number of common shares #	Amount \$
Issued and outstanding		
Balance - December 31, 2018	45,106,401	90,152
Issued for cash consideration, net of issuance costs	5,404,855	26,957
Stock options exercised	105,196	353
Warrants exercised	14,423	82
Balance - December 31, 2019	<u>50,630,875</u>	<u>117,544</u>
Issued for cash, net of issuance costs	274,790	345
Stock options exercised	77,264	239
DSUs redeemed	76,920	184
Balance - March 31, 2020	<u>51,059,849</u>	<u>118,312</u>

As at March 31, 2020, a total of 2,009,874 shares (2019 - 2,069,142) are reserved to meet outstanding stock options and DSUs.

On March 17, 2020, the Corporation entered into an Equity Distribution Agreement (“ATM Distribution”) with Piper Sandler & Co. (“Piper Sandler”) authorizing the Corporation to offer and sell common shares from time-to-time up to an aggregate offering amount of US\$30,000 through Piper Sandler, as agent. Management estimates that the total expenses associated with the ATM Distribution, excluding compensation and reimbursements payable to Piper Sandler under the terms of the Equity Distribution Agreement, will be approximately \$426. As of March 31, 2020, 274,790 common shares have been sold under the ATM Distribution for total gross proceeds of \$771. An additional 624,620 common shares were sold subsequent to March 31, 2020 for gross proceeds of \$1,753, bringing total proceeds raised under the Equity Distribution Agreement to \$2,524 as of May 14, 2020.

On March 6, 2019, the Corporation completed a public offering, issuing an aggregate of 4,900,000 common shares at a price of \$5.45 per common share, raising gross proceeds of \$26,705. On March 11, 2019, the underwriters partially exercised their option to purchase common shares, resulting in the issuance of 504,855 common shares of the Corporation at a price of \$5.45 per share for additional gross proceeds of approximately \$2,751. As a result of the exercise of this option, the Corporation raised total gross proceeds of approximately \$29,456 before deducting the underwriting commissions and offering expenses of \$2,499.

IMV Inc.

Notes to the Unaudited Interim Condensed Consolidated Financial Statements

As at March 31, 2020 and December 31, 2019

(Expressed in thousands of Canadian dollars except for share and per share amounts)

6 Contributed surplus

	Amount
	\$
Contributed surplus	
Balance - December 31, 2018	6,504
Share-based compensation:	
Stock options vested	1,138
DSUs vested	290
Reclassification of DSUs	955
Stock options exercised	(258)
Warrants expired	62
Balance - December 31, 2019	<u>8,691</u>
Share-based compensation:	
Stock options vested	306
DSUs vested	122
Stock options exercised	(206)
DSUs Redeemed	(189)
Warrants expired	332
Balance - March 31, 2020	<u><u>9,056</u></u>

Deferred share units

The maximum number of common shares which the Corporation is entitled to issue from Treasury in connection with the redemption of DSUs granted under the DSU Plan is 468,750 common shares.

DSU activity for the three months ended March 31, 2020 and the year ended December 31, 2019 are as follows:

	March 31,	December 31,
	2020	2019
	#	#
Opening balance	360,965	223,604
Granted	50,422	137,361
Redeemed	(79,106)	-
Closing balance	<u><u>332,281</u></u>	<u><u>360,965</u></u>

At March 31, 2020, there were 332,281 (2019 - 360,965) DSUs outstanding related to this Plan.

Compensation expense at March 31, 2020 was \$122 (2019 - (\$191) recovery), recognized over the vesting period. All services received in exchange for the grant of DSUs were measured at their fair values at the time of grant and vest immediately. Vested DSUs cannot be redeemed until the holder is no longer a member of the Board.

On August 8, 2019 (the "Reclassification Date"), the the Corporation resolved to settle all future DSU redemptions in shares, instead of cash. All outstanding DSUs are accordingly now considered equity-settled instruments. As a result of this change, the fair value of the DSUs at the Reclassification Date were reclassified from liabilities to contributed surplus.

IMV Inc.

Notes to the Unaudited Interim Condensed Consolidated Financial Statements

As at March 31, 2020 and December 31, 2019

(Expressed in thousands of Canadian dollars except for share and per share amounts)

6 Contributed surplus (continued)**Stock options**

The fair values of stock options are estimated using the Black-Scholes option pricing model. As at March 31, 2020, 245,850 stock options (2019 - 343,100) with a weighted average exercise price of \$5.98 (2019 - \$6.39) and a term of five years (2019 - five years), were granted to employees and consultants. The expected volatility of these stock options was determined using historical volatility rates and the expected life was determined using the weighted average life of past options issued. The value of these stock options has been estimated at \$757 (2019 - \$1,112), which is a weighted average grant date value per option of \$3.08 (2019 - \$3.24), using the Black-Scholes valuation model and the following weighted average assumptions:

	2020	2019
Risk-free interest rate	1.38%	1.81%
Expected volatility	66%	64%
Expected life (years)	4.2	4.2
Forfeiture rate	4%	5%
Expected dividend yield	0%	0%

Option activity for the three months ended March 31, 2020 and the year ended December 31, 2019 was as follows:

	March 31, 2020		December 31, 2019	
	Number	Weighted average exercise price	Number	Weighted average exercise price
	#	\$	#	\$
Outstanding - Beginning of year	1,573,411	4.63	1,474,477	4.12
Granted	245,850	5.98	343,100	6.39
Exercised	(115,626) ¹	2.16	(139,877) ¹	2.32
Forfeited	-	-	(90,570)	6.81
Cancelled	(26,042)	6.40	(1,219)	7.04
Expired	-	-	(12,500)	2.37
Outstanding - End of year	1,677,593	4.96	1,573,411	4.63

¹ Of the 115,626 (2019 - 139,877) options exercised, 100,001 (2019 - 98,408) elected the cashless exercise, under which 61,639 shares (2019 - 63,727) were issued. These options would have otherwise been exercisable for proceeds of \$217 (2019 - \$229) on the exercise date.

The weighted average exercise price of options exercisable at March 31, 2020 is \$4.17 (2019 - \$3.29).

The maximum number of common shares issuable under the Corporation's stock option plan shall not exceed 4,600,000, inclusive of all shares presently reserved for issuance pursuant to previously granted stock options.

IMV Inc.

Notes to the Unaudited Interim Condensed Consolidated Financial Statements

As at March 31, 2020 and December 31, 2019

(Expressed in thousands of Canadian dollars except for share and per share amounts)

7 Warrants

Warrant activity for the period ended March 31, 2020 and the year ended December 31, 2019 was as follows:

	March 31, 2020			December 31, 2019		
	Number #	Weighted average exercise price	Amount	Number #	Weighted average exercise price	Amount
		\$	\$		\$	\$
Opening balance	134,766	6.53	332	192,458	5.84	415
Exercised	-	-	-	(14,423)	4.22	(21)
Expired	(134,766)	6.53	(332)	(43,269)	4.22	(62)
Closing balance	-	-	-	134,766	6.53	332

The fair values of warrants are estimated using the Black-Scholes option pricing model. There were no warrants issued in 2019 or 2020.

8 Financial instruments**Fair value of financial instruments**

Financial instruments are defined as a contractual right or obligation to receive or deliver cash on another financial asset.

The following table sets out the approximate fair values of financial instruments as at the unaudited interim condensed consolidated statements of financial position date with relevant comparatives:

	March 31, 2020		December 31, 2019	
	Carrying value	Fair value	Carrying value	Fair value
	\$	\$	\$	\$
Cash and cash equivalents	7,372	7,372	14,066	14,066
Amounts receivable	318	318	439	439
Accounts payable and accrued liabilities	7,470	7,470	6,142	6,142
Amounts due to directors	64	64	60	60
Long-term debt	8,871	8,871	8,461	8,461

Assets and liabilities, such as commodity taxes, that are not contractual and that arise as a result of statutory requirements imposed by governments, do not meet the definition of financial assets or financial liabilities and are, therefore, excluded from amounts receivable and accounts payable and accrued liabilities.

Fair value of items, which are short-term in nature, have been deemed to approximate their carrying value. The above noted fair values, presented for information only, reflect conditions that existed only at March 31, 2020 and December 31, 2019 and do not necessarily reflect future value or amounts which the Corporation might receive if it were to sell some or all of its assets to a willing buyer in a free and open market.

IMV Inc.

Notes to the Unaudited Interim Condensed Consolidated Financial Statements

As at March 31, 2020 and December 31, 2019

(Expressed in thousands of Canadian dollars except for share and per share amounts)

8 Financial instruments (continued)

The fair value of the long-term debt is estimated based on the expected interest rates for similar borrowings by the Corporation at the unaudited interim condensed consolidated statements of financial position dates. At March 31, 2020, the fair value is estimated to be equal to the carrying amount.

9 Related party transactions

During the three months ended March 31, 2020, there were no related party transactions (2019 - \$nil).

10 Subsequent events

On May 7, 2020, the Corporation completed a non-brokered private placement, issuing an aggregate of 8,770,005 Units at the market price of \$2.86 per Unit, raising gross proceeds of \$25,100. Each Unit will consist of one common share of the Corporation and 0.35 of one common share purchase warrant ("Warrant"). Each Warrant will have an exercise price of \$3.72 and will be exercisable until 24 months after its issuance. Management estimates that the total expenses associated with the private placement, will be approximately \$125.

(9)



Management's Report on Financial Position and Operating Results

For the three months ended March 31, 2020

LETTER TO SHAREHOLDERS

Dear Fellow Shareholders,

Amidst these very challenging times created by COVID-19, IMV continues to leverage the versatility of its platform to produce cancer-targeted immunotherapies and vaccines against infectious diseases that program immune cells in vivo. Every day we strive to deliver this novel class of immunotherapies and vaccines, applying the unique mechanism of action of our DPX technology to elicit targeted, robust and sustained immune responses against important human diseases with high unmet medical needs.

In response to the pandemic, we implemented measures in line with recommended public health interventions to ensure the safety of our team and have been in active communication with clinical sites to preserve patient safety and continuity of critical care in our trials. As we navigate these challenges, we are reminded daily of our work's importance and remain committed to our mission.

Over the last few months, we have continued to validate our DPX platform and advance our clinical pipeline. In particular, we are pleased to share an important update from SPiReL, a Phase 2 study of a DPX-Survivac combination regimen in patients with recurrent/refractory diffuse large B-cell lymphoma (r/r DLBCL). At the American Society for Hematology (ASH) Annual Meeting in December 2019, we shared preliminary results from this study: 56% of evaluable subjects exhibited clinical responses, including three complete responses (3/9) and two partial responses (2/9). Today, we are very pleased to report that this study, which will enroll up to 25 patients, has already met its primary endpoint with seven clinical responses in the first eleven evaluable patients or a clinical response rate of 64%. This represents a significant milestone for DPX-Survivac as we look forward sharing detailed results from this study at a scientific conference later in 2020.

Among other recent accomplishments from our lead program were promising interim Phase 2 results from DeCidE1, in which DPX-Survivac achieved sustained clinical activity while being very well tolerated in patients with advanced and recurrent ovarian cancer. Notably, these results showed a 53% tumor regression rate and a 79% disease control rate in a heavily pre-treated population and compared favorably to the existing standard of care against multiple key metrics. Taken together, these results underscore DPX-Survivac's ability to shrink both solid and hematological tumors, with long-lasting clinical responses and a differentiated safety profile.

Although our focus remains in oncology, we are also pursuing other opportunities with partners to leverage our DPX technology for other serious diseases. In recognition of the global public health crisis surrounding COVID-19, we joined forces with some of the most recognized vaccine researchers and clinicians in Canada to tackle this disease. Moving rapidly since identifying target epitopes in late March, we have already successfully manufactured and formulated preclinical batches of DPX-COVID-19 based on these targets. With partners, IMV has also been conducting preclinical studies and generating data to support the selection of the most promising epitope candidates to advance into human clinical studies. As part of this process, we are currently finalizing our Phase 1 design and have submitted to relevant authorities requests for grants to fund our DPX-COVID-19 program.

From a financial perspective, we recently completed a private placement, delivering gross proceeds of \$25.1M CAD from notable new and existing institutional investors. This financing strengthens our financial position and based on our current plan provides funds to support clinical development of DPX-Survivac, research and development activities and other corporate costs through year-end 2021.

Despite these challenging times, we are looking ahead to the remainder of 2020 with optimism. At the upcoming ASCO Virtual Scientific Program which will be held at the end of May, we expect to report translational and updated clinical response data from the DeCidE1 trial of DPX-Survivac in advanced and recurrent ovarian cancer patients. Additionally, we look forward to reporting topline Phase 2 results from the DPX-Survivac basket trial and SPiReL study in r/r DLBCL before year-end. Beyond our lead program, we are preparing to advance our clinical candidate for COVID-19 as quickly as possible and continue to leverage our platform against other targets of interest.

As illustrated above, we continue to make great progress in unlocking the value of our DPX platform for patients afflicted with cancer and other serious diseases, including COVID-19. We are grateful for the continued support of our partners, shareholders and employees and we look forward working closely with them as we continue to deliver on IMV's great opportunities.

/s/ Frederic Ors

Frederic Ors
Chief Executive Officer

MANAGEMENT DISCUSSION AND ANALYSIS (“MD&A”)

The following analysis provides a review of the unaudited interim condensed consolidated results of operations, financial condition, and cash flows for the three months ended March 31, 2020 (“Q1 2020”), with information compared to the three months ended March 31, 2019 (“Q1 2019”), for IMV Inc. (“IMV or the Corporation”). This analysis should also be read in conjunction with the information contained in the audited consolidated financial statements and related notes for the years ended December 31, 2019 and December 31, 2018.

The Corporation prepares its unaudited interim condensed consolidated financial statements in accordance with International Financial Reporting Standards (“IFRS”) as issued by the International Accounting Standards Board (IASB). Management is responsible for the preparation of the consolidated financial statements and other financial information relating to the Corporation included in this report. The Board of Directors is responsible for ensuring that management fulfills its responsibilities for financial reporting. In furtherance of the foregoing, the Board of Directors has appointed an Audit Committee comprised of independent directors. The Audit Committee meets with management and the auditors in order to discuss the results of operations and the financial condition of the Corporation prior to making recommendations and submitting the consolidated financial statements to the Board of Directors for its consideration and approval for issuance to shareholders. The information included in this MD&A is as of May 14, 2020, the date when the Board of Directors approved the Corporation’s unaudited interim condensed consolidated financial statements for the three months ended March 31, 2020, on the recommendation of the Audit Committee.

Amounts presented in this MD&A are approximate and have been rounded to the nearest thousand except for per share data. Unless specified otherwise, all amounts are presented in Canadian dollars.

Additional information regarding the business of the Corporation, including the Annual Information Form of the Corporation for the year ended December 31, 2019 (the “AIF”) and included in the Corporation’s registration statement on Form 40-F filed with the U.S. Securities and Exchange Commission, is available on SEDAR at www.sedar.com and on EDGAR at www.sec.gov/edgar.

FORWARD-LOOKING STATEMENTS

Certain statements in this MD&A may constitute “forward-looking” statements which involve known and unknown risks, uncertainties and other factors that may cause the actual results, performance, or achievements of the Corporation, or industry results, to be materially different from any future results, performance, or achievements expressed or implied by such forward-looking statements. When used in this MD&A, such statements use such words as “will”, “may”, “could”, “intends”, “potential”, “plans”, “believes”, “expects”, “projects”, “estimates”, “anticipates”, “continue”, “potential”, “predicts” or “should” and other similar terminology. These statements reflect current expectations of management regarding future events and operating performance and speak only as of the date of this MD&A. Forward-looking statements include, among others:

- The Corporation’s business strategy;
- Statements with respect to the sufficiency of the Corporation’s financial resources to support its activities;
- Potential sources of funding;
- The Corporation’s ability to obtain necessary funding on favorable terms or at all;
- The Corporation’s expected expenditures and accumulated deficit level;
- The Corporation’s expected outcomes from its ongoing and future research and research collaborations;
- The Corporation’s ability to obtain necessary regulatory approvals;
- The Corporation’s expected outcomes from its pre-clinical studies and trials;
- The Corporation’s exploration of opportunities to maximize shareholder value as part of the ordinary course of its business through collaborations, strategic partnerships, and other transactions with third parties;
- The Corporation’s plans for the research and development of certain product candidates;
- The Corporation’s strategy for protecting its intellectual property;
- The Corporation’s ability to identify licensable products or research suitable for licensing and commercialization;
- The Corporation’s ability to obtain licences on commercially reasonable terms;
- The Corporation’s plans for generating revenue;
- The Corporation’s plans for future clinical trials; and

- The Corporation's hiring and retention of skilled staff.

Forward-looking statements involve significant risks and uncertainties, should not be read as guarantees of future performance or results, and will not necessarily be accurate indications of whether or not such results will be achieved. A number of factors could cause actual results to differ materially from the results discussed in the forward-looking statements, including, but not limited to, the factors discussed in the AIF, under the heading "Risk Factors and Uncertainties". Although the forward-looking statements contained in this MD&A are based upon what management of the Corporation believes are reasonable assumptions, the Corporation cannot provide any assurance to investors that actual results will be consistent with these forward-looking statements and should not be unduly relied upon by investors.

Actual results, performance and achievements are likely to differ, and may differ materially, from those expressed or implied by the forward-looking statements contained in this MD&A. Such statements are based on a number of assumptions which may prove to be incorrect, including, but not limited to, assumptions about:

- Obtaining additional funding on reasonable terms when necessary;
- Positive results of pre-clinical studies and clinical trials;
- The Corporation's ability to successfully develop existing and new products;
- The Corporation's ability to hire and retain skilled staff;
- The products and technology offered by the Corporation's competitors;
- General business and economic conditions, including as a result of the pandemic outbreak of COVID-19;
- The Corporation's ability to protect its intellectual property;
- The Corporation's ability to manufacture its products and to meet demand;
- The general regulatory environment in which the Corporation operates; and
- Obtaining necessary regulatory approvals and the timing in respect thereof.

These statements reflect management's current views and beliefs and are based on estimates, assumptions, and information currently available to, and considered reasonable by, management. The forward-looking information in this MD&A does not include a full assessment or reflection of the unprecedented impacts of the COVID-19 pandemic occurring in the first half of 2020 and the ongoing and developing resulting indirect global and regional economic impacts. The Corporation is currently experiencing uncertainty related to the rapidly developing COVID-19 situation. Uncertainties include the scope, severity and duration of the pandemic, the actions taken to contain or mitigate its impact and the direct and indirect effect of the pandemic and containment measures, among others. It is anticipated that the spread of COVID-19 and global measures to contain it will have an impact on the Corporation, however it is challenging to quantify the potential magnitude of such impact at this time. The Corporation is regularly assessing the situation and remains in contact with its partners, clinical sites and investigators, and suppliers to assess any impacts and risks.

The information contained herein is dated as of May 14, 2020, the date of the Board's approval of the Q1 2020 unaudited interim condensed consolidated financial statements and of the MD&A. For additional information on risks, uncertainties, and assumptions, including a more detailed assessment of the risks that could cause actual results to materially differ from current expectations, please refer to the AIF of IMV filed on SEDAR at www.sedar.com and included in the registration statement on Form 40-F filed on EDGAR at www.sec.gov/edgar.

CORPORATE OVERVIEW

IMV is a clinical-stage biopharmaceutical company dedicated to making immunotherapy more effective, more broadly applicable, and more widely available to people facing cancer, infectious and other serious diseases. IMV is pioneering a new class of immunotherapies based on the Corporation's proprietary drug delivery platform ("**DPX**"). This patented technology leverages a novel mechanism of action ("**MOA**") discovered by the Corporation. This MOA does not release the active ingredients at the site of injection but forces an active uptake by immune cells (antigen-presenting cells) and delivery of active ingredients into lymph nodes. This unique MOA enables the programming of immune cells *in vivo*, which are aimed at generating powerful target-specific therapeutic capabilities. DPX's no-release MOA can be leveraged to generate "first-in-class" T cell therapies with the potential, in the opinion of IMV, to be disruptive in the treatment of cancer.

The Corporation's first cancer immunotherapy uses survivin-based peptides licensed from Merck KGaA, on a world-wide exclusive basis, formulated in DPX (**DPX-Survivac**). Survivin is a well characterized and tumor-associated antigen known to be overexpressed in more than 20 different cancers. DPX-Survivac leverages the MOA of the DPX platform to generate a constant flow of killer T cells in the blood that are targeted against survivin expressed on cancer cells. It is comprised of five minimal major histocompatibility complex ("**MHC**") class I peptides to activate naïve T cells against survivin.

DPX-Survivac is currently being tested in:

- A phase 2 clinical trial that evaluates DPX-Survivac in an open label safety and efficacy study in ovarian cancer patients with advanced platinum-sensitive and resistant ovarian cancer;
- Two investigator-sponsored phase 2 clinical trials in combination with the checkpoint inhibitor Keytruda® (pembrolizumab) of Merck & Co Inc. (“**Merck**”) in patients with recurrent, platinum-resistant and sensitive ovarian cancer and in patients with measurable or recurrent diffuse large B cell lymphoma (“**DLBCL**”); and
- A phase 2 basket trial in combination with Merck’s Keytruda® (pembrolizumab) in patients with select advanced or recurrent solid tumors in bladder, liver (hepatocellular carcinoma), ovarian, or non-small-cell lung (“**NSCLC**”) cancers, as well as tumors shown to be positive for the microsatellite instability high (“**MSI-H**”) biomarker.

In infectious disease indications, the Corporation has completed a demonstration phase 1 clinical trial with a target against the respiratory syncytial virus (“**RSV**”). The Corporation also has a commercial licensing agreement with Zoetis for the development of two targeted therapies for cattle and is also conducting several research and clinical collaborations, including a collaboration with the Dana-Farber Cancer Institute (“**Dana-Farber**”) for Human Papillomavirus (“**HPV**”) related cancers and with Leidos, Inc. (“**Leidos**”) in the United States for the development of targeted therapies for malaria and the Zika virus.

The Corporation is also developing a vaccine candidate against COVID-19 (“**DPX-COVID-19**”) in collaboration with lead investigators for the phase 1 clinical study: Joanne Langley, M.D. and Scott Halperin, M.D., of the Canadian Center for Vaccinology (“**CCIV**”) at Dalhousie University, the Izaak Walton Killam Health Center and the Nova Scotia Health Authority and the Canadian Immunization Research Network (“**CIRN**”); along with Dr. Gary Kobinger, Ph.D., Director of the Research Centre on Infectious Diseases at the University Laval in Quebec City and Global Urgent and Advanced Research and Development (“**GUARD**”) in Canada.

The common shares of the Corporation (the “**Common Shares**”) are listed on the Nasdaq Stock Market LLC (“**Nasdaq**”) and on the Toronto Stock Exchange (“**TSX**”) under the symbol “**IMV**”.

BUSINESS MODEL AND STRATEGY

IMV is dedicated to making immunotherapy more effective, more broadly applicable, and more widely available to people facing cancer and other serious diseases. The Corporation’s lead product candidate, DPX-Survivac, has demonstrated the ability to induce prolonged T cell activation leading to tumor regressions in advanced ovarian cancer and DLBCL.

Foremost, the Corporation’s clinical strategy is to target late stage unmet medical needs for a shorter path to clinical demonstration and first regulatory approval. In addition, the Corporation is evaluating combination with Merck’s Keytruda® checkpoint inhibitor in multiple solid tumor indications.

In collaboration with commercial and academic partners, the Corporation is also expanding the application of DPX as a delivery platform for other applications. Pre-clinical and clinical studies have indicated to date that the Corporation’s delivery platform may allow for the development of enhanced targeted therapies for a wide range of infectious diseases by generating a stronger and more durable immune response than with existing delivery methods.

The Corporation intends to be opportunistic in the development of products by exploring a variety of avenues, including co-development through potential collaborations, strategic partnerships or other transactions with third parties. The Corporation may seek additional equity and non-dilutive funding and partnerships to advance the development of its product candidates.

PLATFORM AND PRODUCTS IN DEVELOPMENT

Delivery Platform

The DPX platform is a unique and patented formulation discovered by the Corporation that provides a new way to deliver active ingredients to the immune system using a novel MOA. This MOA does not release the active ingredients at the site of injection but forces an active uptake by immune cells (antigen-presenting cells) and delivery of active ingredients into lymph nodes. IMV is exploiting this unique MOA to pioneer a new class of immunotherapies that represents a paradigm shift from current approaches. Thanks to its “no release” MOA, the DPX-based targeted therapies allow the programming of immune cells *in-vivo* to generate new target-specific therapeutic capabilities. The DPX platform can be leveraged to generate “first-in-class” T cell therapies with the potential to be disruptive in the treatment of cancer. The Corporation believes that the novel MOA of DPX makes the platform uniquely suitable for cancer immunotherapies, which are designed to target tumor cells. DPX-based candidates can induce prolonged, target-specific, and polyfunctional T cell activation, which are postulated to be required for effective tumor control.

The DPX platform is based on active ingredients formulated in lipid nanoparticles and, after freeze drying, suspended directly into a lipidic formulation. DPX-based products are stored in a dry format, which provides the added benefit of an extended shelf life. The formulation is designed to be easy to re-suspend and administer to patients.

DPX also has multiple manufacturing advantages: it is fully synthetic; can accommodate hydrophilic and hydrophobic compounds; is amenable to a wide-range of applications (for example, peptides, small-molecules, RNA/DNA, or antibodies); and provides long term stability as well as low cost of goods.

The DPX platform forms the basis of all IMV's product development programs.

DPX-Survivac

Product Candidate Overview

DPX-Survivac, the Corporation's first cancer immunotherapy candidate, uses survivin-based peptides licensed from Merck KGaA on a world-wide exclusive basis that are formulated in DPX. DPX-Survivac leverages the MOA of DPX to generate a constant flow of T cells in the blood that are targeted against survivin expressed on cancer cells and is comprised of five minimal MHC class I peptides to activate patients' naïve T cells against survivin.

Survivin is a well characterized and recognized tumor associated antigen known to be expressed during fetal development and across most tumor cell types, but it is rarely present in normal non-malignant adult cells. Survivin controls key cancer processes (apoptosis, cell division, and metastasis) and has been associated with chemoresistance and cancer progression. It has been shown that survivin was expressed in all 60 different human tumor lines used in the National Cancer Institute's cancer drug screening program and is documented in the literature to be overexpressed in more than 20 indications.

Cancer	Survivin %
Ovarian	90
Breast	90
Melanoma	90
Lung	53
Colorectal	54
Gastric	94
Kidney	23-82
Glioblastoma	80
ALL	70
CML	70
MDS	90
DLBCL	80

Figure 1: Examples of % of patients with survivin expression in different indications

In clinical trials exploring the activity of DPX-Survivac, an intermittent low-dose oral regimen of cyclophosphamide is used as an immune-modulator. Conventional chemotherapeutic drugs are traditionally used for their cytotoxic effect on tumors but cyclophosphamide can also be used at lower doses to potentiate the activity of other immunotherapies without inducing significant cytotoxicity.

Several studies have demonstrated that low-dose regimens of cyclophosphamide can have multiple beneficial effects for T cell therapies such as DPX-Survivac, including reduction of T regulatory cell numbers and increase in effector T cells (Hugues et al, Immunology, 2018). In Phase 1 clinical studies, IMV demonstrated that intermittent low-dose oral cyclophosphamide can act as an immune-modulator increasing the number of survivin-specific T cells generated by DPX-Survivac (Weir et al, AACR, 2016).

CLINICAL PIPELINE

	Product (target)	Indication	Preclinical	Phase 1	Phase 2	Phase 3	Sponsor	Collaborators
Immunotherapies	DPX-Survivac/CPA (Survivin)	Ovarian	▶				IMV™	
		DLBCL	▶				Sunnybrook RESEARCH INSTITUTE	MERCK
		Basket Trial: Lung (NSCLC), Bladder, Liver, Ovarian, MSI-H	▶				IMV™	MERCK
	DPX-SurMAGE /CPA (Survivin + MAGE A9)	Bladder	▶			IMV™	CHU de Québec Université Laval	
	DPX-BRAF/CPA (BRAF)	Melanoma	▶			IMV™	THE WESTAR INSTITUTE	
Vaccines	DPX-RSV (SheA)	Respiratory Syncytial Virus (RSV)	▶				IMV™	CIRN
	DPX-COVID-19 (Spike)	COVID-19	▶				IMV™	CIRN

IMMUNO-ONCOLOGY

DPX-Survivac is being tested in 6 different cancer indications through multiple phase 2 clinical trials.

DPX-Survivac - Ongoing Clinical Trials

COVID-19 Impact on Clinical Program

It is anticipated that the COVID-19 pandemic crisis will impact ongoing trial activities across the industry due to the pressure placed on the healthcare system as well as governmental and institutional restrictions. IMV’s clinical team is working closely with each clinical site and our contract research organizations (“CRO”)s on a contingency plan to ensure that patient safety and the integrity of data is maintained. IMV is following the FDA guidance issued for the COVID-19 pandemic: “FDA Guidance on Conduct of Clinical Trials of Medical Products during COVID-19 Pandemic Guidance for Industry, Investigators, and Institutional Review Boards”. Additionally, the team continues to monitor updated institutional, regional and national guidance to fully comply with applicable guidelines as they are issued. It is noted that some clinical sites have paused or slowed enrollment in clinical trials, while other sites, less impacted, are continuing activities as planned. The overall enrollment rate may decrease, but clinical activities are continuing. Patients are encouraged to comply with directives from public health officials and, subject to such compliance, attend visits as planned or to discuss alternatives with their physician. The current activities performed at central labs to assess the eligibility of patients and the management of clinical samples is not impacted to date, and IMV is working with the vendors to ensure continuity of activities. Drug supply is not expected to be impacted at this time. As added precaution, IMV is working on a contingency plan to ensure proper provisioning of drugs to all clinical sites in the event of future transportation or other constraints.

Ovarian subpopulation - DeCidE1 phase 1b/2

The DeCidE1 phase 2 study is a multicenter, randomized, open-label study to evaluate the safety and effectiveness of DPX-Survivac with intermittent low dose cyclophosphamide (CPA). This phase 2 arm enrolled 22 patients with recurrent, advanced platinum-sensitive and resistant ovarian cancer. Patients received 2 subcutaneous injections of DPX-Survivac 3 weeks apart and every eight weeks thereafter, and intermittent low dose CPA one week on and one week off for up to 1 year. Paired tumor biopsies were performed prior to treatment and on treatment.

Primary endpoints of this study are overall response rate, disease control rate and safety. Secondary endpoints include cell mediated immunity, immune cell infiltration in paired biopsy samples, duration of response, time to progression, overall survival and biomarker analyses.

On February 4, 2020, the Corporation presented clinical translational data supporting the mechanism of action of its lead compound, DPX-Survivac/CPA during the 2020 ASCO-SITC Clinical Immuno-Oncology Symposium. The Corporation measured systemic immune responses, tumor immune infiltrates and clinical tumor response from pre- and post-treatment patient samples in connection with three Phase 1 and/or Phase 2 clinical studies, each evaluating DPX-Survivac/CPA alone or in a combination regimen in patients with platinum-sensitive or resistant, advanced ovarian cancer. Highlights from these translational data include:

- DPX-Survivac generates robust, functional, targeted, and sustained survivin-specific T cell response in ovarian cancer subjects in the maintenance setting as well as with recurrent disease.
- DPX-Survivac induced activation of cytolytic T cell pathway is correlated with clinical response highlighting its unique mechanism of action.
- Enhanced number of unique survivin-specific T cell clones are detected in on-treatment tumor samples and the T cell infiltration on-treatment correlated with clinical responses.
- DPX-Survivac mechanism of action has been confirmed across multiple clinical trials and has shown to provide clinical benefit and long-term clinical response in some subjects with advanced recurrent ovarian cancer.

On February 25, 2020, the Corporation reported updated results from the ongoing DeCidE1 Phase 2 study of DPX-Survivac/CPA, in patients with advanced recurrent ovarian cancer. The new results show that DPX-Survivac immunotherapy is active and well-tolerated.

19 patients were evaluable for efficacy with six patients (31%) still receiving treatment. Key preliminary findings are outlined below:

- 15 patients (79%) achieved disease control, defined as Stable Disease (“SD”) or Partial Response (“PR”) on target lesions:
 - Tumor shrinkage of target lesions was observed in 10 patients (53%).
- Durable clinical benefits lasting ≥ 6 months were observed in seven patients (37%) so far:
 - Four of these seven patients (21% of evaluable patients) achieved PR with tumor regression $>30\%$ on target lesions;
 - Three stable diseases were ongoing for > 6 months (range 7-9) including -29.5% and -12% tumor regressions; and
 - Median duration not reached yet, with five of these seven (71%) patients still on treatment at > 6 months (range 7-10).
- Analysis of Baseline Tumor Burden (“BTB”) showed durable clinical benefits across a broad range of BTB (1.5-7.7 cm) with a higher number of patients achieving benefits in BTB < 5 cm as previously observed in other arms of the study:
 - Six out 11 with BTB < 5 cm (55%) achieved clinical benefits lasting > 6 months.
- Durable clinical benefits include platinum-resistant and refractory patients who previously received PARP inhibitors and bevacizumab.
- Treatment was well-tolerated, with most adverse events being Grade 1-2 reactions at the injection site.

IMV plans to take these results to the U.S. Food and Drug Administration (“FDA”) for a Type B meeting, to align on the design of a Phase 2b study with potential to support registration under accelerated approval in this indication.

Translational and updated clinical response data from Phase 2 DeCidE1 study of DPX-Survivac in advanced ovarian cancer to be presented by Dr. Oliver Dorigo, MD, Ph.D., at the upcoming 2020 ASCO 2020 Virtual Scientific Program which will be held from May 29th to May 31st.

In December 2018, IMV met with the FDA in a Type B meeting to discuss the results to date of its DeCidE1 clinical trial and continuing development plan, as well as to obtain agency guidance on a potential accelerated regulatory pathway for DPX-Survivac as a T cell immunotherapy for the treatment of advanced ovarian cancer in patients with progressing disease.

The purpose of IMV's Type B meeting with the FDA was to request feedback on the design of the clinical program for DPX-Survivac. This program includes the continuing DeCidE1 phase 2 clinical study and a potential future registration trial for accelerated approval in a subset of ovarian cancer patients.

The FDA reviewed the Corporation's proposed clinical development plan and acknowledged the potential for accelerated approval in advanced ovarian cancer based on objective response rate ("ORR") according to Recist 1.1 criteria with reported median duration of response ("DOR"). In addition, the FDA provided important guidance on clinical design considerations for different lines of therapy and platinum-sensitive and resistant patient populations.

Drug	Registrational Clinical Trials	Indication	Base for approval
Olaparib (Lynparza) Approved December 2014	Single arm, open label, Phase 2 (Study 42)	Germline BRCA mutation ≥3 prior lines of chemotherapy	N=137 ORR: 34% - platinum-resistant: 30% mDoR: 7.9 mo
Rubraca (Rucaparib) Breakthrough in April 2015 Approved December 2016	Single arm, open label, Phase 2 (Study 10 and ARIEL2)	Germline and/or somatic BRCA mutation ≥2 prior lines of chemotherapy	N=106 ORR: 42% platinum-resistant: 25% mDoR: 6.7 mo-9.2 mo

Figure 2: Examples of previous US FDA accelerated approvals in ovarian cancer (source: FDA website)

In addition, IMV submitted a protocol amendment for a predictive enrichment approach to the phase 2 DeCidE1 trial, and further discussed those details with the FDA during the Type B meeting. The phase 2 primary end point, based on ORR per Recist 1.1 criteria, is intended to confirm the high response rate and duration of clinical benefits observed in previously announced results in a patient population defined by a clinical biomarker based on baseline tumor burden.

The Corporation believes that there is still an urgent medical need in advanced recurrent ovarian cancer (Sources: 1. NCCN Guidelines Ovarian Cancer V2.2018; SEER Ovarian Cancer; JCO, vol 33; 32 Nov 2015, Gyn Onc 133(2014) 624-631):

- Nearly 70% of ovarian cancers are diagnosed in advanced stage;
- The overall 5-year survival rate is 46.5%, and only 29% for advanced disease;
- Most patients develop advanced, platinum-resistant, poor prognosis disease; and
- Limited options exist with current single-agents at 6-30% response rates and median progression free survival ("mPFS") of 2.1 - 4.2 months.

The Corporation believes that it has the potential to be "best-in-class" in the competitive landscape of recurrent ovarian cancer as other immunotherapeutic treatments tested in this patient population (Merck's Keytruda, and Pfizer/Merck KGaA's Bavencio) are unlikely to proceed into registration trials based on the published results available:



Figure 3: Recurrent ovarian cancer immunotherapy competitive landscape

The Corporation's clinical strategy with this trial is to establish the targeted T cell activity of its lead compound in order to increase value and de-risk clinical development, and to target late stage unmet medical needs for a shorter path to clinical demonstration and first regulatory approval.

The Corporation anticipates that, in addition to general clinical expenses which are distributed amongst the various clinical projects, the costs to complete this phase 2 clinical trial is currently estimated at \$750,000 which is expected to be spent in 2020.

Combinations with Merck's Keytruda® (pembrolizumab)

Phase 2 clinical trial in DLBCL - SPiReL Phase 2 (investigator-sponsored)

This phase 2 study is a combination trial with Merck's Keytruda® (pembrolizumab) in patients with measurable or recurrent DLBCL led by Dr. Neil Berinstein, MD, FRCP(c), ABIM, hematologist-oncologist at the Odette Cancer Centre at Sunnybrook Health Sciences Centre (investigator-sponsored). This investigator sponsored trial, announced initially in May 2017, is designed to evaluate the safety and efficacy of DPX-Survivac, Merck's Keytruda® (pembrolizumab), and intermittent low-dose cyclophosphamide. IMV has provided an update on this trial at the American Society of Hematology Annual meeting held on December 6-10, 2019.

The primary objective of this study is to document the response rate to this treatment combination using modified Chesonⁱ criteria. Secondary objectives include duration of response and safety. Exploratory endpoints include T cell response, tumor immune cell infiltration, and gene expression analysis.

ⁱ Cheson, B.D., Pfistner, B., Juweid, M.E., Gascoyne, R.D., Specht, L., Horning, S.J. and Diehl, V. (2007). Revised Response Criteria for Malignant Lymphoma. *Journal of Clinical Oncology*, 25(5) DOI: 10.1200/JCO.2006.09.2403.

CR: Nodal disease less than 1.5 cm, absence of extranodal disease, no new lesions and normal bone marrow (BM);

PR: ≥50% decrease in the sum of the product of the diameters (SPD), no new lesion;

PD: Longest diameter of node > 1.5 cm and ≥50% increase from Product of Perpendicular Diameter and increase in longest or smallest diameter from nadir (lowest value), unequivocal progression of non target, new lesions or BM involvement.

This study has met its primary efficacy endpoint with 64% (7/11) of evaluable patients demonstrating a clinical response so far. The study remains ongoing and the top line data are expected to be presented at a conference later in 2020. As of May 7, 2020, 20 subjects have been enrolled across five different clinical sites in Canada.

Researchers conducting the investigator sponsored study are testing the novel immunotherapy combination in patients whose DLBCL expresses survivin, a tumor antigen highly expressed in 60 percent of DLBCL patients. DPX Survivac stimulates the immune system to produce T cell responses targeting survivin.

On December 8, 2019, IMV provided updated data on this study. Seven of the nine evaluable patients demonstrated clinical benefit, including three complete responses and two partial responses.

Updated SPiReL data highlights:

At the time of data cut-off for this analysis, efficacy data based on modified Cheson criteria was available from nine evaluable patients:

- 7/9 (77.8%) evaluable subjects exhibited clinical benefit, including three (33.3%) complete responses and two (22.2%) partial responses;
- Reproducible survivin-specific T cell responses observed in all subjects that achieved clinical responses on treatment;
- One subject, who received three prior lines of systemic therapies and failed autologous stem cell transplant, reached a complete response at the first on-study scan following treatment with the DPX-Survivac combination regimen and remains free of disease recurrence after completing the study; and
- Clinical benefits and favorable toxicity profile observed in a heterogenous population of r/r DLBCL patients, including patients of advanced age and/or with comorbidities, who are more susceptible to adverse effects and more difficult to treat.

The Corporation anticipates that, in addition to general clinical expenses which are distributed amongst the various clinical projects, its share of the cost to complete this study is currently estimated at \$600,000 which is expected to be spent in 2020.

Phase 2 basket trial in 5 solid tumor indications

In September 2018, the Corporation announced the expansion of its clinical program with a phase 2 basket trial in collaboration with Merck evaluating its lead candidate, DPX-Survivac/CPA, and Merck's KEYTRUDA® (pembrolizumab), in patients with select advanced or recurrent solid tumors.

The open-label, multicenter, phase 2 basket study will evaluate the safety and efficacy of the immunotherapeutic combination in patients with bladder, liver (hepatocellular carcinoma), ovarian, or non-small cell lung NSCLC cancers, as well as tumors shown to be positive for the microsatellite instability high (MSI-H) biomarker. Investigators plan to enroll up to 184 patients across five indications in 20 medical centers in Canada and the United States.

The ASCO defines a basket clinical study as a trial that investigates the effects of a drug regimen in multiple tumor types that share a common molecular target, regardless of where the disease originated.

This is the third clinical trial evaluating the combination of DPX-Survivac/CPA and KEYTRUDA® (pembrolizumab) in advanced recurrent cancers.

On September 30, 2019, IMV presented preliminary results from its ongoing phase 2 basket trial, during the Immunotherapy of Cancer poster session at the European Society for Medical Oncology (ESMO) 2019 Congress in Barcelona, Spain.

Preliminary Results from the Phase 2 Basket Trial

At the time of cut-off, 23 patients were enrolled across all five patient cohorts. This includes 19 patients across all cohorts who received DPX-Survivac in combination with pembrolizumab with CPA, and four patients from the ovarian cancer cohort receiving DPX-Survivac with only pembrolizumab:

- Preliminary results from the first on-study scan showed tumor reduction in patients with ovarian cancer (with and without CPA), NSCLC and bladder cancer;
- Partial responses observed at first scan in two subjects (bladder cancer, ovarian cancer); 19/23 subjects are still active on study treatment;
- T cell infiltration observed in biopsy samples from subjects who achieved tumor reduction on treatment;
- Eight ovarian cancer patients were enrolled in the study, randomized 1:1 to treatment with and without CPA. Tumor control and tumor reductions were observed in both groups; and
- Safety evaluation on all evaluable patients demonstrated that treatment was well-tolerated, with no related Grade 3-4 or immune-related adverse events (AEs) reported.

As at May 7, 2020, 19 clinical sites were open, and 92 patients had been enrolled across the five indications. Given the ongoing COVID-19 pandemic, the pressure placed on the healthcare system, as well as governmental and institutional restrictions, the enrollment in this trial as well as the collection and analysis of data has been slowed down.

IMV is uncertain of when it will be able to report results from this trial but expects to disclose interim data in the second half of 2020 when a more mature dataset will be available. The Corporation anticipates that, in addition to general clinical expenses, which are distributed amongst the various clinical projects, \$22,400,000 is currently estimated to be spent for stage 1 for this trial, of which \$6,500,000 is estimated to be spent in 2020.

Phase 2 clinical trial in ovarian cancer (investigator-sponsored)

In February 2017, the Corporation announced an investigator-sponsored phase 2 clinical trial in ovarian cancer in combination with Merck's checkpoint inhibitor pembrolizumab in patients with recurrent, platinum-resistant ovarian cancer. University Health Network's ("UHN") Princess Margaret Cancer Centre conducts the phase 2 non-randomized, open-label trial designed to evaluate the potential anti-tumor activity of the combination of pembrolizumab, DPX-Survivac, and intermittent low-dose cyclophosphamide. It is expected to enroll 42 subjects with advanced epithelial ovarian, fallopian tube, or primary peritoneal

cancer. The study's primary objective is to assess overall response rate. Secondary study objectives include progression free survival rate, overall survival rate, and potential side effects, over a five-year period. At this stage, the Corporation has no specific plan on the next steps after this trial as it will have to be assessed with its partner based on the clinical trial results.

As of May 7, 2020, 16 patients were enrolled in the trial and the Corporation will disclose results once provided by the UHN Princess Margaret Cancer Centre and currently anticipates that, in addition to general clinical expenses, which are distributed amongst the various clinical projects, its share of the costs to complete this study, currently expected to be spent in 2020, are estimated at \$200,000.

DPX-SurMAGE

In March 2019, IMV announced that CQDM, a Canadian bioresearch consortium, had awarded a grant for a collaboration among IMV Inc., Centre de recherche du CHU de Quebec-Universite Laval ("CHU") and La Fondation du CHU de Quebec ("FCHUQc"). The collaboration will receive a grant of up to \$1,200 from the CQDM and \$300 from the FCHUQc over three years, to develop a novel dual target T cell therapy for an initial clinical application in bladder cancer. IMV currently expects to contribute \$2,800,000 over the next three years towards this project of which \$1,600,000 has been contributed in 2019 and \$500,000 is estimated to be contributed in 2020.

The work will target immunogenic peptides from the MAGE protein family member A9 (MAGE-A9). This protein is frequently expressed in various human cancers including bladder, lung and kidney. These peptides will be combined with selected immunogenic peptides from the survivin protein composing the DPX-Survivac T cell drug candidate.

The researchers believe that MAGE-A9 and survivin peptides presented on the surface of cancer cells can be used to program T cells to destroy tumors and may represent ideal targets for anti-cancer T cell immunotherapies. The collaborators will combine these peptides with IMV's proprietary DPX technology to develop a first-in-class dual target T cell therapy (DPX-SurMAGE).

DPX-SurMAGE will be initially evaluated in preclinical studies. Upon successful completion of these preclinical evaluations, researchers are aiming to test the candidate in two clinical studies in patients with:

- Muscle invasive bladder cancer combined with an anti-PD-1 and intermittent CPA prior to cystectomy; and
- Low-grade highly recurrent non-muscle invasive bladder cancer combined with CPA prior to transurethral resection.

This collaboration is expected to span a three-year period and as part of the collaboration agreement, IMV holds an exclusive option to in-license intellectual property related to this collaboration.

In June 2019, IMV met with Health Canada for a pre-clinical trial application ("CTA") meeting. The objectives of this meeting were to present and discuss the strategy for the development (including pre-clinical and clinical plans) of DPX-SurMAGE, to the agency to ensure the strategy was aligned with the agency's expectations. The agency agreed with the approach for pre-clinical, manufacturing and clinical development and made suggestions to facilitate its review by the agency.

Given the ongoing COVID-19 pandemic, the pressure placed on the healthcare system, as well as governmental and institutional restrictions, and the fact that IMV had not initiated a phase 1 trial of DPX-SurMAGE prior to the pandemic, IMV is uncertain of when it will initiate this trial. The Corporation intends to provide an update when more information is available.

Orphan Drug Status and Fast Track Designation

The Corporation announced, in November 2016, that the European Medicines Agency (EMA) had granted orphan drug designation status to IMV's DPX-Survivac in ovarian cancer. In July 2015, the FDA also granted orphan drug status to DPX-Survivac for the treatment of ovarian cancer. This designation is valid for all applications of DPX-Survivac in ovarian cancer without restriction to a specific stage of disease.

IMV had previously received FDA fast track designation for DPX-Survivac. The designation is intended for patients with no measurable disease after their initial surgery and chemotherapy.

Other Programs

Oncology

DPX-NEO

On January 17, 2019, treatment of the first patient occurred in the phase 1 trial evaluating neoepitopes formulated in the Corporation's proprietary DPX delivery platform in patients with ovarian cancer. The study is part of the Corporation's DPX-NEO program, which is a continuing collaboration between UConn Health and IMV to develop neoepitope-based anti-cancer therapies.

Investigators will assess the safety and efficacy of using patient-specific neoepitopes discovered at UConn Health and formulated in IMV's proprietary DPX-based delivery technology in women with ovarian cancer. Investigators plan to enroll up to 15 patients in the phase 1 study. UConn Health is financing the trial with IMV providing materials and advice.

The Corporation expects to disclose results only when those are made available by UConn Health.

DPX-E7

Dana-Farber is leading the DPX -E7 study through a \$1.5 million research grant from Stand Up To Cancer and the Farrah Fawcett Foundation to clinically evaluate collaborative translational research that addresses critical problems in HPV-related cancers. The Dana-Farber study is a single center, open label, non-randomized clinical trial that will investigate the safety and clinical efficacy in a total of 44 treated participants. Its primary objectives are to evaluate changes in CD8+ T cells in peripheral blood and tumor tissue, and to evaluate the safety in HLA-A2 positive patients with incurable HPV-related head and neck, cervical, or anal cancers. The trial has pre-consented 76 patients so far, from which 11 patients have been treated.

The Corporation expects to disclose results only when those are made available by Dana-Farber.

Other Applications

Product Overview

A component of the Corporation's business strategy is partnering the DPX platform for infectious and other disease applications. The DPX platform has the potential to generate a rapid and robust immune response, often in a single dose. The unique single-dose capability could prove to be beneficial in targeting difficult infectious and other disease candidates.

DPX-COVID-19

In early March, IMV announced plans to develop a DPX-based vaccine candidate for COVID-19 in collaboration with experts in the field. IMV is on schedule to complete preclinical studies before the end of May and is on track to initiate a Phase 1 clinical study for DPX-COVID-19 this summer.

The ongoing pandemic outbreak of COVID-19 and its alarmingly quick transmission to over 125 countries across the world resulted in the World Health Organization ("WHO") declaring a pandemic on March 11, 2020.

The outbreak is caused by a novel coronavirus, the Severe Acute Respiratory Syndrome Coronavirus 2 ("SARS-CoV-2"). There is an urgent need to develop vaccines to control its spread and help protect vulnerable populations. However, the bottleneck with current conventional vaccine approaches is the length of time required for vaccine development. The Corporation believes IMV's DPX delivery technology offers the possibility of a fully synthetic epitope-based approach with the potential for accelerated development and rapid, large-scale production of a vaccine that would be compliant with current good manufacturing practice ("cGMP").

Research in coronaviruses has identified the benefit of humoral and cellular (B and T cell) immune responses for treatment and protection from infection.

IMV believes that it has already demonstrated in multiple clinical trials in oncology and infectious diseases the potential of its technology for the induction of robust and sustained B and T cells. The Corporation believes there is an opportunity to pursue a COVID-19 development program to establish the clinical safety and immunogenicity using a similar approach for COVID-19.

The Corporation is developing its vaccine candidate DPX-COVID-19 in collaboration with lead investigators for the phase 1 clinical study: Joanne Langley, M.D. and Scott Halperin, M.D., of CCFV at Dalhousie University, the Izaak Walton Killam Health Center and the Nova Scotia Health Authority and CIRN; along with Dr. Gary Kobinger, Ph.D., Director of the Research Centre on Infectious Diseases at the University Laval in Quebec City and GUARD in Canada. The investigators are assisting with preclinical and clinical evaluation and with further development strategy in collaboration with the Canadian government and others.

Since IMV announced its plans on March 18, 2020 to develop a DPX-based vaccine for COVID-19 in collaboration with well-respected experts the Corporation has made significant progress:

- The Corporation has used sequences of the virus and immunoinformatics to predict and identify several hundred epitopes, of which 23 were selected for their biological relevance to the virus and potential to generate neutralizing antibodies against SARS-CoV-2;
- Based on this analysis, peptide candidates targeting these epitopes were manufactured and IMV's suppliers and contract manufacturers were engaged to prepare for the cGMP batch required to support a clinical study in humans;
- In collaboration with Gary Kobinger, PhD, Director of the Research Centre on Infectious Diseases at the University Laval in Quebec City and GUARD in Canada, preclinical assays in animal models are ongoing and will be completed before the end of May to validate the safety and potency of the vaccine candidate before initiating the human clinical study;
- In collaboration with Joanne Langley, MD, at the CCFV and the CIRN, the design of a phase 1 clinical study is in the process of being completed and clinical sites have been identified in both Nova Scotia and Quebec;
- IMV recently held discussions with Health Canada about the design of a phase 1 clinical study in humans scheduled to be initiated this summer. The proposed trial will enroll approximately 84 patients. It will be a randomized, placebo-controlled study that will assess two different doses of DPX-COVID-19 in two age strata; and
- The Corporation has submitted several grant applications in Canada and the United States to help support its COVID-19 clinical program.

Third-party research in related coronaviruses has identified the benefit of humoral and cellular (B and T cell) immune responses for protection and resolution of infection, and the Corporation believes the body of data it has produced to date supports its DPX platform for peptide-based induction of B cells and T cells. The Corporation is now designing a vaccine candidate against COVID-19 based on third-party immunological studies of SARS-CoV and third-party sequencing data available for SARS-CoV-2 with the goal of selecting potentially immunogenic epitopes within the virus that induce neutralizing antibody responses and protective T cell responses.

Through the Corporation's other clinical studies, the Corporation believes its DPX technology has demonstrated a favorable safety profile and immunogenicity in both cancer and infectious disease settings, with sustained effect and potential for single-dose effectiveness as a prophylactic vaccine. Over 200 patients have been dosed with DPX-based immunotherapies and data from these studies suggest treatment is well-tolerated, including in heavily pre-treated cancer patients with advanced-stage disease. The Corporation has also applied this technology for the prevention of RSV, the second-leading cause of respiratory illness in infants, the elderly and the immunosuppressed. The Corporation reported its Phase 1 data from its clinical candidate, DPX-RSV, which demonstrated a favorable safety profile and immunogenicity in older adults (age 50-64), as well as preclinical data from research-stage candidates aimed at other infectious diseases, including malaria and anthrax.

RSV

The Corporation has performed preclinical research activities for an RSV targeted candidate, which is the second leading cause of respiratory illness in infants, the elderly, and the immunosuppressed. Currently, there is no preventive therapy available for this virus and IMV is seeking to develop a novel DPX-based formulation to be used in elderly and healthy adults, including women of child-bearing age. IMV has in-licensed the RSV antigen exclusively from VIB VZW, a non-profit life sciences

research institute funded by the Flemish government, to expand its pipeline of DPX-based candidates. The novel RSV antigen being evaluated in the DPX platform is based on the short hydrophobic protein present at low levels on the surface of the RSV virion. But, more importantly, it is also present on the surface of RSV-infected cells. This DPX-based candidate has a unique mechanism of action in which the resultant antibodies bind to and destroy infected cells.

Phase 1 clinical trial in RSV

A phase 1 clinical study has been conducted in Canada with the Corporation's RSV targeted candidate in healthy adults. The RSV candidate is formulated in IMV's proprietary DPX platform and is initially being developed to protect the elderly population from infection. The phase 1 study, which was the first clinical trial of a DPX-based formulation in an infectious disease indication, evaluated the safety and immune response profile of the DPX-RSV candidate in 40 healthy older adult volunteers (age 50-64 years) and two dose cohorts, with 20 subjects in each cohort.

In October 2016 and April 2017, the Corporation announced positive topline results from this trial. The report outlined that more than nine months after the last vaccination, 15 of 16 participants (93%) who received DPX-RSV demonstrated antigen-specific immune responses. The candidate also continued to have a positive safety profile and was well tolerated with no SAEs among all study participants. Within the 25µg dose patient cohort, which was the only dose tested out to one year, 100 percent of older adults (7/7 immune responders) vaccinated with DPX-RSV maintained the antigen-specific immune responses one year after receiving the booster dose. After one year, the antibody levels measured were still at peak with no sign of decrease.

On September 27, 2018, IMV announced results of ongoing research to further explore the novel MOA of its candidate. New data from a preclinical study highlighted the effects of two potential approaches to preventing RSV, comparing a single dose bovine version of DPX-RSV to a two-dose conventional investigational bovine RSV (bRSV) preventive therapy. Researchers found that IMV's targeted therapy yielded strong antigen-specific immune responses and a protective effect on disease pathology.

They found SH antibodies in 14 of the 15 animals that received DPX-bRSV, and the improvements observed in disease pathology were comparable between the two cohorts. These were the first bovine animal health data to directly correlate the induced immune response against IMV's novel RSV target - the SH viral protein- with measures of disease protection.

Conventional RSV preventive therapies target either the F or G proteins of the virus and provide protection by neutralizing the RSV virus. Clinical measures of efficacy focus on the amount of neutralizing antibodies in the bloodstream. DPX-RSV works differently; it targets the SH viral ectodomain of the RSV virus and, instead of neutralizing the virus, it enables the immune system to recognize and destroy infected cells. Because there are no neutralizing antibodies resulting from the DPX-RSV MOA, a different clinical assessment is required to determine the candidate's protective effect. IMV has exclusive worldwide licenses on applications that target the SH ectodomain antigen in RSV. The Corporation is exploring opportunities to out-license this product to potential partners.

Leidos Collaboration

In 2016, IMV was awarded a subcontract by Leidos, a health, national security, and infrastructure solutions company, to evaluate IMV's DPX platform for the development of peptide-based malaria targets. The subcontract is funded through Leidos' prime contract from the U.S. Agency for International Development ("USAID") to provide DPX-based candidate evaluations in the preclinical, clinical, and field stages of malaria preventative therapy development. Leidos and IMV are working together to identify adjuvant and antigen combinations that can be used to protect against malaria and, with the DPX delivery system, formulate promising targeted therapy candidates for potential clinical testing.

In November 2017, an expansion of this collaboration was announced. Following the achievement of several preclinical milestones in the collaboration with USAID, Leidos and USAID selected the DPX-based platform as one of the preferred formulations for further development under a new contract extension. Under the new subcontract, the collaborators are conducting additional research that focuses on identifying the most promising target-formulation combinations.

Zoetis Collaboration

On August 31, 2017, the Corporation announced the achievement of several milestones in its ongoing collaboration with global animal health company Zoetis to develop targeted T cell therapy for cattle. In recent controlled studies, the IMV

formulations met efficacy and duration of immunity endpoints against two disease targets. These results will enable Zoetis to advance two DPX-formulation candidates into late-stage testing.

Licensing Agreements

While the Corporation is focused on developing a pipeline of cancer immunotherapies, it is also pursuing opportunities to license its platform technology to other parties interested in creating enhanced T cell targeted therapies on an application-by-application basis.

In April 2018, IMV signed a licensing agreement and granted SpayVac-for-Wildlife (SFW Inc.) a license to two of its proprietary delivery platforms. SFW Inc. has global exclusive rights to use both of these platforms to develop humane, immune-contraceptive compounds for control of overabundant, feral and invasive wildlife populations against royalties on sales.

MARKET OVERVIEW

Cancer Immunotherapies

Cancer is considered one of the most widespread and prevalent diseases globally. According to the 2019 Cancer Facts & Figures released by the American Cancer Society, it is predicted that the global cancer burden will rise to 27.5 million and the number of cancer deaths to 16.3 million by 2040 solely due to the growth of the aging population. However, these projections may be underestimates given the adoption of unhealthy behaviors and lifestyles associated with rapid income growth and changes in reproductive patterns in economically transitioning countries. According to the 2019 Cancer Facts & Figures, cancer usually develops in older people; 80% of all cancers in the United States are diagnosed in people 55 years of age or older. The “oldest old”, adults ages 85 and older are the fastest-growing population group in the US and women outnumber men in this age group because of a longer life expectancy.

Conventional cancer treatment involves surgery to remove the tumor whenever possible, as well as chemotherapy and radiation. Chemotherapies are widely used, despite their associated toxicities, because they interfere with the ability of cancer cells to grow and spread. However, studies have shown that older patients often receive little or no treatment because the benefit of prolonged survival does not outweigh potential adverse effects and impact on quality of life. Also, in all groups of patients, tumors often develop resistance to chemotherapies, thus limiting their efficacy in preventing tumor recurrence. Despite recent advances, independent sources note a high unmet medical need in cancer therapy, noting the median survival rate remains poor. Cancer immunotherapies may provide new and effective treatments. According to a Market & Markets report released in September 2016, the global immunotherapy drug market is projected to reach USD\$119.39 billion by 2021 from USD\$61.97 billion in 2016, growing at a compound annual growth rate (“CAGR”) of 14 % during the forecast period of 2016 to 2021. The major players operating in the immunotherapy drug market include F. Hoffmann-La Roche AG (Switzerland), GlaxoSmithKline (U.K.), AbbVie, Inc. (U.S.), Amgen, Inc. (U.S.), Merck (U.S.), Bristol-Myers Squibb (U.S.), Novartis International AG (Switzerland), Eli Lilly and Corporation (U.S.), Johnson & Johnson (U.S.), and AstraZeneca plc (U.K.).

Cancer immunotherapy seeks to harness the immune system to assist in the destruction of tumors and to prevent their recurrence. There has been significant interest in the field of cancer immunotherapy stemming from recent clinical success in prolonging patient survival with novel compounds. The ability to apply these appropriately has resulted from a greater understanding of the immune dysfunction that is characteristic of cancer. One area in which there have been breakthroughs has been in the area of checkpoint inhibitors, which are compounds that target key regulatory molecules of the immune system. Yervoy® (anti CTLA 4, or ipilimumab, developed by Bristol Myers Squibb) was the first compound in this class to be approved for use in advanced metastatic melanoma. In cancer, these regulators (CTLA 4, PD 1 and its ligand PD L1) act to inhibit CD8 T cell-mediated anti-tumor immune responses that are crucial for tumor control. Monoclonal antibodies that target PD 1 and PD L1 have shown unusual efficacy in cancer patients, with a significant percentage of patients experiencing durable response to these therapies. Several of these compounds have been approved in multiple indications. Merck’s Keytruda® (pembrolizumab) and Bristol Myers Squibb’s Opdivo® (nivolumab) received FDA approval in 2014 for advanced melanoma patients who have stopped responding to other therapies. These therapies have subsequently been approved for use in other advanced cancers including bladder cancer, non-small cell lung cancer, Hodgkin’s Lymphoma, squamous cell carcinoma of the head and neck and stomach cancer. In addition, Keytruda® in particular has been approved for use in cancers with a specific molecular indication irrelevant of cancer type, having been approved in May 2017 for use to treat solid tumors having a biomarker for microsatellite instability (MSI-H), which is a defect in the DNA repair pathway. This represents about 5% of a number of different tumor types, including colorectal, breast, prostate, and thyroid cancers.

These drugs have been shown to be helpful in treating several types of cancer but with success only in a limited percentage of patients. It is not yet known exactly why, though researchers have noticed that these drugs seem to work especially well for patients whose cancer cells have a higher number of mutations.

Key opinion leaders in the field have indicated that the solution lies in combining checkpoint inhibitors with other cancer treatments and that the ideal combination is likely to be a therapy that drives tumor specific immune responses. These include novel T cell-based therapies. These targeted therapies fit well with checkpoint inhibition therapy because they simultaneously activate strong tumor-specific T cell activation, while also releasing the brakes on immune suppression. The success of such combinations should allow pharmaceutical companies to significantly expand the market of their checkpoint inhibitors.

The Corporation believes that targeted T cell therapies will become an important component of these novel combination immunotherapies, with the potential of synergistic benefits to become an essential part of a multi-pronged approach for the treatment of cancer.

INTELLECTUAL PROPERTY

The Corporation strives to protect its intellectual property in established, as well as emerging, markets around the world. The Corporation's intellectual property portfolio relating to its platform technology includes 17 patent families, the first of which contains eight patents issued in five jurisdictions (United States, Europe, Canada, Japan, and Australia). The 16 other families collectively contain 41 patents issued in 10 jurisdictions (United States, Europe, Canada, Australia, Japan, India, Israel, Singapore, China, and, separately, Hong Kong) and 61 pending patent applications in 9 jurisdictions. Taking into account the validations of the European patents, the Corporation's intellectual property portfolio includes 94 patents. More details on the Corporation's intellectual property strategy and patents can be found in the AIF filed on SEDAR at www.sedar.com.

The Corporation owns registered trademarks in the United States, Canada, and Europe.

RECENT AND QUARTERLY DEVELOPMENTS

In March 2020, the World Health Organization declared the COVID-19 outbreak a global pandemic. IMV continues to monitor the COVID-19 situation, which is rapidly developing. IMV has been designated as an essential business by the Nova Scotia Department of Business and Nova Scotia Public Health. In addition to adhering to directives from public health officials, IMV has implemented a pandemic contingency plan to guide employees, contractors, visitors, facilities and operations. The Corporation's plan includes identifying essential business activities to help ensure continuity of business, restricting access to its offices and operation sites and encouraging all employees to work from home to the extent possible, asking business partners to engage by telephone or video conference where possible, eliminating business travel and requiring self-isolation for employees travelling outside of Canada and increasing the frequency and emphasis on cleaning and sanitizing. As the COVID-19 health-crisis further develops, IMV will continue to rely on guidance and recommendations from local health authorities and the Centers for Disease Control and Prevention to update its policies.

The Corporation announced:

- On May 7, 2020, the completion of a private placement (the "Private Placement") of 8,770,005 units of the Company (each, a "Unit") at the market price of \$2.86 per Unit. With aggregate gross proceeds of approximately \$25.1 million this non-brokered private placement is being co-led by Fonds de Solidarité FTQ, an existing investor, and Lumira Ventures, a new investor in the Company, along with participation by Altium Capital, also a new investor in IMV, together with incumbent investors.
- On March 30, 2020, that it has made significant progress on the development of DPX-COVID-19, a vaccine candidate against the novel coronavirus, including:
 - The Corporation has used sequences of the virus and immunoinformatics to predict and identify several hundred epitopes, of which 23 were selected for their biological relevance to the virus and potential to generate neutralizing antibodies against SARS-CoV-2;
 - Based on this analysis, IMV has begun manufacturing peptide candidates targeting these epitopes as well as planning with IMV's suppliers and contract manufacturers to prepare for the cGMP batch required to support a clinical study in humans;

- In collaboration with Gary Kobinger, Ph.D., Director of the Research Centre on Infectious Diseases at the University Laval in Quebec City, preclinical assays in animal models are also planned in April through May of this year to validate the safety and potency of the vaccine candidate before initiating the human clinical study;
- In collaboration with Joanne Langley, M.D. at the Canadian Center for Vaccinology (CCfV) and the Canadian Immunization Research Network (CIRN) the design of a Phase 1 clinical study in 48 healthy subjects has been completed and clinical sites identified in both Nova Scotia and Quebec;
- IMV has initiated discussions with Health Canada in preparation for a CTA. A meeting is being scheduled in the week of April 20, 2020 with the goal to initiate the clinical study in the summer of 2020; and
- The Corporation has submitted several grant applications in Canada in an effort to help support its clinical program.
- On March 18, 2020, that it is advancing the clinical development of a DPX-based vaccine candidate against COVID-19. The goal of the development program, in collaboration with lead investigators for the phase 1 clinical study: Joanne Langley, M.D. and Scott Halperin, M.D., of the CCfV at Dalhousie University, the Izaak Walton Killam Health Center and the Nova Scotia Health Authority and the CIRN; along with Dr. Gary Kobinger, Ph.D., Director of the Research Centre on Infectious Diseases at the University Laval in Quebec City and GUARD in Canada, is to establish the clinical safety and immunogenicity of a vaccine candidate based on the Corporation's DPX delivery technology and incorporating peptides targeting novel epitopes from the coronavirus strain.
- On March 18, 2020, that it has entered into an equity distribution agreement with Piper Sandler & Co. ("Piper Sandler") pursuant to which the Corporation may, from time to time sell, through "at-the-market offerings" with Piper Sandler acting as sales agent, on the Nasdaq such number of Common Shares as would have an aggregate offering price of up to US\$30 million (the "ATM Distribution"). The Corporation plans to use the net proceeds from the ATM Distribution, if any, for general corporate purposes, including but not limited to working capital expenditures, capital expenditures, research and development expenditures, and clinical trial expenditures, including expenditures related to a COVID-19 vaccine candidate.
- On February 25, 2020, that updated results from DeCidE1, an ongoing Phase 2 study of its lead candidate, DPX-Survivac, in patients with advanced recurrent ovarian cancer were reported during a conference call and webcast.

All 22 patients with advanced recurrent ovarian cancer enrolled in this arm of the study were heavily pre-treated, with the median number of prior therapies greater than three.

As of February 24, 2020, 19 patients were evaluable for efficacy with six patients (31%) still receiving treatment. Key preliminary findings are outlined below:

- 15 patients (79%) achieved disease control, defined as Stable Disease (SD) or Partial Response (PR) on target lesions:
 - Tumor shrinkage of target lesions was observed in 10 patients (53%).
- Durable clinical benefits lasting ≥ 6 months were observed in seven patients (37%) so far:
 - Four of these seven patients (21% of evaluable patients) achieved PR with tumor regression $>30\%$ on target lesions;
 - Three stable diseases were ongoing for > 6 months (range 7-9) including -29.5% and -12% tumor regressions; and
 - Median duration not reached yet, with five of these seven (71%) patients still on treatment at > 6 months (range 7-10).
- Analysis of Baseline Tumor Burden (BTB) showed durable clinical benefits across a broad range of BTB (1.5-7.7 cm) with a higher number of patients achieving benefits in BTB < 5 cm as previously observed in other arms of the study:
 - Six out 11 with BTB < 5 cm (55%) achieved clinical benefits lasting > 6 months.
- Durable clinical benefits include platinum-resistant and refractory patients who previously received PARP inhibitors and bevacizumab; and
- Treatment was well-tolerated, with most adverse events being Grade 1-2 reactions at the injection site.

- On February 14, 2020, that Albert Scardino was to retire from the IMV Board of Directors effective February 28, 2020.
- On February 4, 2020, the presentation of clinical translational data supporting the mechanism of action of its lead compound, DPX-Survivac, during the 2020 ASCO-SITC Clinical Immuno-Oncology Symposium, being held in Orlando, FL.

As part of this analysis, the Corporation measured systemic immune responses, tumor immune infiltrates and clinical tumor response from pre- and post-treatment patient samples in connection with three Phase 1 and/or Phase 2 clinical studies, each evaluating DPX-Survivac alone or in a combination regimen in patients with platinum-sensitive or resistant, advanced ovarian cancer. Highlights from these translational data include:

- DPX-Survivac generated survivin-specific T cells in the blood of 80% of patients sampled;
- Clinical anti-tumor responses were correlated with increased infiltration of T cells into tumors following treatment with DPX-Survivac;
- DPX-Survivac induced enrichment in T cell, cytotoxic lymphocytes and B cell-specific signatures which correlate with clinical response; and
- Antigen-specific T cells retained their functionality throughout the duration of treatment.

SELECTED FINANCIAL INFORMATION

The selected statements of loss and comprehensive loss data for the periods presented and the selected statement of financial position data as of the dates presented are derived from the unaudited interim condensed consolidated financial statements. The selected historical financial data below should be read in conjunction with the financial statements and related notes and the sections titled "Components of Operations Overview" and "Results of Operations" appearing elsewhere in this report.

	Three months ended March 31,	
	2020	2019
Statements of loss and comprehensive loss data:		
(in thousands, except share and per share amounts)		
Revenue		
Subcontract revenue	\$ -	\$ 8
Interest revenue	68	74
Total revenue	68	82
Operating Expenses		
Research and development	6,824	4,013
General and administrative	3,033	1,960
Government assistance	(558)	(346)
Accreted interest	433	398
Total operating expenses	9,732	6,025
Net loss and comprehensive loss	\$ (9,664)	\$ (5,943)
Basic and diluted loss per share	\$ (0.19)	\$ (0.13)
Weighted-average shares outstanding	50,719,488	46,712,436

	As of,	
	March 31, 2020	December 31, 2019
Statement of financial position data:		
(in thousands of Canadian dollars)		
Cash and cash equivalents	\$ 7,372	\$ 14,066
Working capital (1)	4,553	13,199
Total assets	15,329	22,434
Total liabilities	17,744	15,986
Accumulated deficit	(129,783)	(120,119)
Total shareholder's equity (deficit)	(2,415)	6,448

(1) Working capital is defined as current assets less current liabilities. See financial statements for further details regarding current assets and current liabilities.

COMPONENTS OF OPERATIONS OVERVIEW

Revenue

The Corporation has no products approved for commercial sale and has not generated any revenue from product sales. Revenue consists primarily of income earned on cash balances held at a commercial bank. The Corporation also generates immaterial revenue from providing formulation services under research collaboration agreement with Leidos for the development of targeted therapies for malaria and the Zika virus. Revenue is recognized when the formulation services are performed.

Operating Expenses

Research and development expenses

To date, the Corporation's research and development expenses have related primarily to discovery efforts and preclinical, manufacturing and clinical development of its product candidates. The most significant research and development expenses for

the year relate to costs incurred for the development of the Corporation's most advanced product candidates, DPX-Survivac and DPX-SurMAGE, which include:

- Expenses incurred under agreements with CROs, as well as investigative sites and consultants that conduct clinical trials, preclinical studies and other scientific development services;
- Costs related to the production and scale-up of clinical materials, including fees paid to contract manufacturers;
- Employee-related expenses, including salaries, related benefits, travel and share-based compensation expense for employees engaged in research and development functions;
- Expenses incurred for outsourced professional scientific and regulatory development services;
- Laboratory materials and supplies used to support research activities; and
- Facilities and other expenses, which includes depreciation on laboratory equipment.

The Corporation expenses all research and development costs in the periods in which they are incurred. The Corporation accrues for costs incurred as the services are being provided by monitoring the status of the project and the invoices received from its external service providers. Accruals are adjusted as actual costs become known. Where contingent milestone payments are due to third parties under research and development arrangements or license agreements, the milestone payment obligations are expensed when the milestone results are achieved.

Research and development activities are central to IMV's business model. Product candidates in later stages of clinical development generally have higher development costs than those in earlier stages of clinical development, primarily due to the increased size and duration of later-staged clinical trials. The Corporation expects that research and development expenses will increase substantially over the next few years as it increases personnel, advances manufacturing processes, initiates and conducts additional clinical trials and prepares regulatory filings related to its product candidates. The Corporation also expects to incur increased research and development expenses as it selectively identifies and develops additional product candidates.

However, it is difficult to determine with certainty the duration and completion costs of current or future preclinical programs and clinical trials of product candidates.

The duration and timing of clinical trials and development of the Corporation's product candidates will depend on a variety of factors that include, but are not limited to, the following:

- The scope, progress, outcome and costs of clinical trials and other research and development activities, including establishing an appropriate safety profile with IND-directed studies;
- Patient enrollment, discontinuation rates, per patient trial costs, and number and location of clinical trial sites in clinical trials;
- The ability of the Corporation's clinical partners and sponsors for investigator-sponsored trials to manage clinical trials;
- Establishing commercial manufacturing capabilities or making arrangements with third party manufacturers;
- Timing, receipt and terms of any marketing approvals from applicable regulatory authorities;
- Obtaining, maintaining, defending and enforcing patent claims and other intellectual property rights;
- Significant and changing government regulation; and
- Significant competition and rapidly changing technologies within the biopharmaceutical industry.

The probability of success for each product candidate is highly uncertain. The Corporation will determine which programs to pursue and what resources to allocate to each program in response to the scientific and clinical success of each product candidate as well as an assessment of each product candidate's commercial potential. Further, because IMV's product candidates are still in clinical development, the Corporation cannot estimate the actual amounts necessary to successfully complete the development and commercialization of product candidates or whether, or when, it may achieve profitability.

General and administrative

General and administrative expenses consist primarily of salaries and other staff-related costs, including share-based compensation expense for personnel in executive, finance, human resources, project management, business development, investor relations and administrative functions. General and administrative expenses also include, but are not limited to, facilities and overhead costs, legal fees related to corporate, securities and patent matters, investor relations costs, insurance and professional fees for assurance, taxation, information technology communications and human resources matters. General and administrative costs are expensed as incurred and the Corporation accrues for services provided by third parties related to the above expenses by monitoring the status of services provided and receiving estimates from its service providers, adjusting accruals as actual costs become known.

The Corporation expects that its general and administration expenses will increase in the future as it increases personnel to support the continued development of its product candidates. The Corporation has experienced and expects to continue to experience, increased expense associated with being a Nasdaq listed company including increased accounting, audit, legal, regulatory and compliance costs, director and officer insurance premiums, as well as higher investor relations and public relations costs.

Government Assistance

Government assistance consists primarily of research and development investment tax credits awarded through the Canada Revenue Agency's Scientific Research and Economic Development ("SR&ED") program for research expenditures incurred in Canada. Government assistance also contains other government funding for research projects and employment funding as well as fair market value adjustments to interest-free and low-interest government loans.

Accreted interest

Accreted interest relates entirely to the valuation of interest-free and low interest-bearing government loans, most of which are repayable based on a percentage of future gross revenue.

RESULTS OF OPERATIONS

Comparison of the Three Months Ended March 31, 2020 and 2019

The following table summarizes the Corporations results of operations for the three months ended March 31, 2020 and 2019 (in thousands of Canadian dollars):

	Three months ended March 31,		Change (\$)
	2020	2019	
Revenue			
Subcontract revenue	\$ -	\$ 8	\$ (8)
Interest revenue	68	74	-6
Total revenue	68	82	-14
Operating Expenses			
Research and development	6,824	4,013	2,811
General and administrative	3,033	1,960	1,073
Government assistance	(558)	(346)	(212)
Accreted interest	433	398	35
Total operating expenses	9,732	6,025	3,707
Net loss and comprehensive loss	<u>\$ (9,664)</u>	<u>\$ (5,943)</u>	<u>\$ (3,721)</u>

Revenue

Revenue did not significantly fluctuate period over period.

Research and development expenses

Research and development expenses increased to \$6.8 million for the three months ended March 31, 2020 from \$4.0 million for the three months ended March 31, 2019. The increase of \$2.8 million is mainly attributable to \$1 million in clinical costs related to the basket trial as a result of a spike in enrollment in late 2019/early 2020, \$1.3 million increase in non-recurring purchases in 2020 of GMP grade raw materials for DPX-Survivac, \$240,000 in preclinical expenses relating to the DPX-SurMAGE collaboration with CQDM, CHU, and FCHUQc, \$125,000 related to pre-clinical expenses for development of DPX-COVID-19, and \$200,000 in personnel costs due to an increase in headcount. The purchase of GMP grade materials for DPX-Survivac in 2019 and Q1 2020 has covered all the needs of the Corporation for ongoing DPX-Survivac trials until mid 2021.

General and administrative expenses

General and administrative expenses increased to \$3 million for the three months ended March 31, 2020 from \$1.96 million for the three months ended March 31, 2019. The increase of \$1 million compared with Q1 2019 can be explained by an increase of \$157,000 in legal and recruiting fees, \$120,000 in conference and roadshow travel, \$132,000 foreign exchange loss, \$246,000 in investor relations consulting fees, and a \$325,000 increase in deferred share unit (“DSU”) compensation compared with Q1 2019. In Q1 2019, DSU compensation was a (\$204,000) recovery due to outstanding DSUs being revalued each period and a lower share price at the end of Q1 2019, compared with Q4 2018. Effective August 8, 2019, the Corporation elected to settle all future DSU redemptions in shares. As a result, DSUs are now accounted for as equity-settled instruments and will not need to be revalued at each reporting period. The Corporation expects that this will reduce the comparative volatility in the DSU compensation expense from Q3 2020 onward.

Government Assistance

The increase in government assistance for the period ended March 31, 2020 compared with March 31, 2019 is mainly attributable to the increase in SR&ED investment tax credits consistent with increased spend on R&D salaries, raw materials as well as increased clinical trial activity being performed in Canada.

CASHFLOWS, LIQUIDITY AND CAPITAL RESOURCES

Liquidity and Capital Resources

The Corporation has incurred losses and negative cash flows from operations since inception. As of March 31, 2020, the Corporation had an accumulated deficit of \$130 million and anticipates that it will continue to incur net losses for the foreseeable future.

At March 31, 2020, the Corporation had approximately \$10 million of existing and identified potential sources of cash including:

- cash and equivalents of \$7.4 million; and
- amounts receivable and investment tax credits receivable of \$2.6 million.

Management believes that its cash resources of \$7.4 million, its additional potential cash resources of \$2.6 million and the \$25.1 million in gross proceeds from the May 2020 Private Placement, will be sufficient to fund operations for more than 12 months based on current forecasts. This estimate does not take into account any utilization of the ATM Distribution allowing the Corporation to offer and sell Common Shares from time-to-time up to an aggregate offering amount of US\$30 million through Piper Sandler, as agent. As of May 14, 2020, 889,410 Common Shares have been sold under the ATM Distribution for total gross proceeds of US\$1.8 million. The Corporation continually reassesses the adequacy of its cash resources, evaluating existing clinical trials, research projects and/or potential collaboration opportunities, to determine when and how much additional funding is required.

The Corporation’s primary use of cash is to fund operating expenses, which consist primarily of funding clinical and preclinical trials, research and development expenditures and related personnel costs and to a lesser extent, general and administrative expenditures. Cash used to fund operating expenses is impacted by the timing of when the Corporation pays these expenses, as reflected in the change in its outstanding accounts payable and accrued expenses.

It is common for early-stage biotechnology companies to require additional funding to further develop product-candidates until successful commercialization of at least one product candidate. The Corporation’s product candidates are still in the early stages

of clinical and preclinical development and the outcome of these efforts is uncertain. Accordingly, the Corporation cannot estimate the actual amounts necessary to successfully complete the development and commercialization of its product candidates or whether, or when, it may achieve profitability. Until such time, if ever, as the Corporation can generate substantial product revenue, it expects to finance cash needs through a combination of equity or debt financings and collaboration arrangements. If the Corporation does raise additional capital through public or private equity offerings, the ownership interest of its existing stockholders will be diluted, and the terms of these securities may include liquidation or other preferences that adversely affect its stockholders' rights. If IMV raises additional capital through debt financing, it may be subject to covenants limiting or restricting its ability to take specific actions, such as incurring additional debt, making capital expenditures or declaring dividends. If the Corporation is unable to raise capital when needed, it will need to delay, reduce or terminate planned activities to reduce costs. Doing so will likely harm the Corporation's ability to execute its business plans. The Corporation continuously monitors its liquidity position, the status of its development programs including those of its partners, cash forecasts for completing various stages of development, the potential to license or co-develop each product candidate, and continues to actively pursue alternatives to raise capital, including the sale of its equity securities, debt and non-dilutive funding.

Cash Flows

The following table summarizes the Corporation's cash flows for the periods indicated (in thousands of Canadian dollars):

	Three months Ended March 31,	
	2020	2019
Net cash (used in) provided by:		
Operating activities	(6,951)	(8,206)
Financing activities	290	27,630
Investing activities	(33)	(112)
Net increase (decrease) in cash and cash equivalents	<u>(6,694)</u>	<u>19,312</u>

Cashflows from operating activities

During Q1 2020, \$6.95 million was used in operating activities. This included the reported net loss of \$9.66 million prior to being decreased by \$1.02 million for non-cash expenses including DSU compensation, depreciation, accretion of long-term debt and lease obligations and stock-based compensation. The Corporation had a net increase of cash of \$1.7 million as a result of changes in working capital balances, which was mainly attributable to a \$1.37 million increase in accounts payable and accrued liabilities, a \$356,000 decrease in prepaid expenses, and a \$192,000 decrease in amounts receivable, partly offset by an increase of \$225,000 in investment tax credits receivable.

During Q1 2019, \$8.21 million was used in operating activities. This included the reported net loss of \$5.94 million prior to being decreased by \$739,000 for non-cash expenses including DSU compensation, depreciation, accretion of long-term debt and lease obligations, loss on disposal of assets and stock-based compensation. The Corporation had a net decrease of cash of \$3.0 million as a result of changes in working capital balances, which was mainly attributable to a \$3.3 million decrease in accounts payable and accrued liabilities, partly offset by a decrease of \$459,000 in amounts receivable.

Cashflows from financing activities

During Q1 2020, sources of cash from financing activities included: \$771,000 proceeds raised from the ATM Distribution less cash issuance costs of \$426,000, and \$33,000 through the exercise of stock options and warrants. The Corporation used \$83,000 to repay long-term debt and lease obligations during the period.

During Q1 2019, sources of cash from financing activities included: \$29.5 million of proceeds raised in the March 2019 Public Offering less cash issuance costs of \$1.9 million; and \$116,000 through the exercise of stock options and warrants. The Corporation used \$80,000 to repay long-term debt and lease obligations during the period.

Cashflows from investing activities

During Q1 2020, IMV used \$33,000 of cash in investing activities, consisting mainly of purchases of furniture and equipment for ongoing research and operating activities.

During Q1 2019, IMV used \$112,000 of cash in investing activities, consisting mainly of purchases of furniture and equipment for ongoing research and operating activities.

JUNE 2017 EQUITY OFFERING AND USE OF PROCEEDS

On June 21, 2017, the Corporation completed a public offering, issuing 2,403,846 Common Shares at a price of \$4.16 per share for aggregate proceeds of \$10 million. The Corporation intended to use the net proceeds of this offering for the research and development and clinical advancement of its cancer and infectious disease therapy candidates and for working capital and general corporate purposes. The table below provides the amount used to date and any variances in thousands of Canadian dollars (except for working capital and general corporate purposes).

Intended Use of Proceeds	Estimated amount \$	Amount to date \$	Variances
phase 2 clinical trial in DLBCL with Merck	2,400	2,027	No variances anticipated
phase 1 clinical trial for multiple indications	4,200	4,200	None

MARCH 2019 EQUITY OFFERING AND USE OF PROCEEDS

On March 6, 2019, the Corporation completed a public offering, issuing 5,404,855 Common Shares (including 504,855 Common Shares upon the exercise of the underwriters' over-allotment option on March 11, 2019) at a price of \$5.45 per share for aggregate proceeds of \$29.5 million. The Corporation intends to use the net proceeds of this offering to accelerate the development of DPX-Survivac in combination with Keytruda as part of the basket trial in selected advanced or recurrent solid tumors in bladder, liver (hepatocellular carcinoma), ovarian and non-small-cell lung cancers, as well as tumors shown to be positive for the microsatellite instability high biomarker and for general corporate purposes. The table below provides the amount used to date and any variances in thousands of Canadian dollars (except for working capital and general corporate purposes).

Intended Use of Proceeds	Estimated amount \$	Amount to date \$	Variances
Phase 2 clinical trial for multiple indications	16,000	2,671	No variances anticipated

MARCH 2020 ATM DISTRIBUTION AND USE OF PROCEEDS

On March 17, 2020, the Corporation entered into an Equity Distribution Agreement with Piper Sandler & Co. ("Piper Sandler") authorizing the Corporation to offer and sell, through "at-the-market" offerings, common shares from time to time up to an aggregate offering price of US\$30 million through Piper Sandler, as agent. The Corporation intends to use the net proceeds from this offering for research and development expenditures, clinical trial expenditures, including expenditures related to a COVID-19 vaccine candidate and general corporate purposes. As at May 14, 2020, 889,410 common shares have been sold under this agreement for total gross proceeds of US\$1.8 million.

SUMMARY OF QUARTERLY RESULTS

The selected quarterly financial information⁽¹⁾ for the past eight financial quarters is outlined below: (in thousands of dollars, except for amounts per share)

	Q1-2020	Q4-2019	Q3-2019	Q2-2019	Q1-2019	Q4-2018	Q3-2018	Q2-2018
Total Revenue	68	136	164	186	82	133	125	129
Total Expenses	9,732	8,611	8,060	5,237	6,025	7,818	6,112	5,325
Loss	(9,664)	(8,475)	(7,896)	(5,051)	(5,943)	(7,685)	(5,987)	(5,196)
Basic and Diluted Loss per Share	(0.19)	(0.17)	(0.16)	(0.10)	(0.13)	(0.17)	(0.14)	(0.12)

(1) Unless otherwise noted, financial information in thousands of Canadian dollars and prepared in accordance with IFRS.

Revenues from quarter-to-quarter may vary significantly. Revenues are non-recurring by nature and are generated by license agreements as well as contract research agreements. It is also important to note that historical patterns of expenses cannot be taken as an indication of future expenses. The amount and timing of expenses and availability of capital resources vary substantially from quarter-to-quarter, depending on the level of R&D activities being undertaken at any time and the availability of funding from investors or collaboration partners.

OUTLOOK FOR 2020

Milestones	Key dates
Update on the Phase 2 clinical results in the ovarian monotherapy trial at ASCO 2020	Q2 2020
Potential initiation of phase 1 clinical trial for DPX-COVID 19	Summer 2020
Top line phase 2 clinical results update in the DLBCL combination trial	H2 2020
Updated phase 2 clinical results for Basket trial	H2 2020

The exact timing could differ from expectations but are currently management's best estimate.

RELATED PARTY TRANSACTIONS

For the period ending March 31, 2020, there were no related party transactions (2019 - \$nil).

CONTRACTUAL OBLIGATIONS

There is no material change in the contractual obligations of the Corporation since the beginning of the 2020 fiscal year. Details on the contractual obligations of the Corporation can be found in the in the annual audited consolidated financial statements and related notes for the year ended December 31, 2019.

OFF-BALANCE SHEET ARRANGEMENTS

The Corporation was not party to any off-balance sheet arrangements as of March 31, 2020.

OUTSTANDING SECURITIES

As at May 14, 2020, the number of issued and outstanding common shares was 60,454,474 and a total of 5,079,375 stock options, warrants and deferred share units were outstanding.

RISKS AND UNCERTAINTIES

The Corporation is a clinical-stage company that operates in an industry that is dependent on a number of factors that include the Corporation's capacity to raise additional funding on reasonable terms when necessary, obtain positive results of pre-clinical studies and clinical, successfully develop existing and new products, to hire and retain skilled staff, protect its intellectual property, manufacture its products and to meet demand, and obtain necessary regulatory approvals and the timing in respect thereof, etc. An investment in the Common Shares is subject to a number of risks and uncertainties. An investor should carefully consider the risks described in the Corporation's AIF and the registration statement on Form 40-F filed with the U.S. Securities and Exchange Commission, as well as the other information filed with the securities regulators before investing in the Common Shares. If any of the such described risks occur, or if others occur, the Corporation's business, operating results and financial condition could be seriously harmed and investors may lose a significant proportion of their investment.

There are important risks which management believes could impact the Corporation's business. For information on risks and uncertainties, please also refer to the "Risk Factors" section of the Corporation's most recent AIF filed on SEDAR at www.sedar.com and included in the registration statement on Form 40-F filed on EDGAR at www.sec.gov/edgar.

DISCLOSURE CONTROLS AND PROCEDURES AND INTERNAL CONTROLS OVER FINANCIAL REPORTING

Disclosure Controls and Procedures

The Chief Executive Officer (the "CEO") and the Chief Financial Officer (the "CFO") of the Corporation are responsible for establishing and maintaining the Corporation's disclosure controls and procedures ("DCP") including adherence to the Disclosure Policy adopted by the Corporation. The Disclosure Policy requires all staff to keep senior management fully apprised of all material information affecting the Corporation so that they may evaluate and discuss this information and determine the appropriateness and timing for public disclosure.

The Corporation maintains DCP designed to ensure that information required to be disclosed in reports filed under applicable securities laws, is recorded, processed, summarized and reported within the appropriate time periods and that such information is accumulated and communicated to the Corporation's management, including the CEO and CFO, to allow for timely decisions regarding required disclosure.

The CEO and CFO have evaluated whether there were changes to the DCP during the period ended March 31, 2020 that have materially affected, or are reasonably likely to materially affect, the DCP. No such changes were identified through their evaluation.

In designing and evaluating DCP, the Corporation recognizes that any disclosure controls and procedures, no matter how well conceived or operated, can only provide reasonable, not absolute, assurance that the objectives of the control system are met, and management is required to exercise its judgment in evaluating the cost-benefit relationship of possible controls and procedures.

Internal Control over Financial Reporting

The Corporation's management, including the CEO and the CFO, are responsible for establishing and maintaining adequate internal control over financial reporting ("ICFR") for the Corporation to provide reasonable assurance regarding the reliability of financial reporting and the preparation of financial statements for external purposes in accordance with IFRS. The fundamental issue is ensuring all transactions are properly authorized and identified and entered into a well-designed, robust and clearly understood accounting system on a timely basis to minimize risk of inaccuracy, failure to fairly reflect transactions, failure to fairly record transactions necessary to present financial statements in accordance with IFRS, unauthorized receipts and expenditures, or the inability to provide assurance that unauthorized acquisitions or dispositions of assets can be detected.

The CEO and CFO have evaluated whether there were changes to the ICFR during the period ended March 31, 2020 that have materially affected, or are reasonably likely to materially affect, the ICFR. No such changes were identified through their evaluation. In response to the COVID-19 pandemic, the Corporation asked its employees to work from home to the extent possible. This change requires certain processes and controls that were previously done or documented manually to be completed and retained in electronic form. Despite the changes required by the current environment, there have been no significant changes in the Corporation's internal controls during the quarter ended March 31, 2020 that have materially affected, or are reasonably likely to materially affect, ICFR. In response to the COVID-19 pandemic, the Corporation asked its employees to work from home to the extent possible. This change requires certain processes and controls that were previously done or documented manually to be completed and retained in electronic form. Despite the changes required by the current environment, there have been no significant changes in the Corporation's internal controls during the quarter ended March 31, 2020 that have materially affected, or are reasonably likely to materially affect, ICFR.

The Corporation's ICFR may not prevent or detect all misstatements because of inherent limitations. Additionally, projections of any evaluation of effectiveness to future periods are subject to the risk that controls may become inadequate because changes in conditions or deterioration in the degree of compliance with the Corporation's policies and procedures.

BASIS OF PRESENTATION OF CONSOLIDATED FINANCIAL STATEMENTS AND SIGNIFICANT ACCOUNTING POLICIES

The consolidated financial statements have been prepared in accordance with the IFRS as issued by the IASB. The accounting policies, methods of computation and presentation applied in the consolidated financial statements are consistent with those of previous financial year except for business development and investor relations expenses are now presented in general and administrative expenses on the consolidated statements of loss and comprehensive loss. Certain comparative figures have been reclassified to conform the presentation adopted in the current year for general and administrative expenses.

The significant accounting policies of IMV are detailed in the notes to the annual audited consolidated financial statements for the year ended December 31, 2019 filed on SEDAR www.sedar.com and included in the registration statement on Form 40-F filed on EDGAR at www.sec.gov/edgar.

CRITICAL ACCOUNTING ESTIMATES AND JUDGEMENTS

Estimates and assumptions are continually evaluated and are based on historical experience and other factors, including expectations of future events that are believed to be reasonable under the circumstances. The determination of estimates requires the exercise of judgement based on various assumptions and other factors such as historical experience and current and expected economic conditions. Actual results could differ from those estimates.

The Corporation's significant accounting policies and critical judgements in applying the Corporation's accounting policies are detailed in the annual audited consolidated financial statements for the year ended December 31, 2019 filed on SEDAR www.sedar.com and included in the registration statement on Form 40-F filed on EDGAR at www.sec.gov/edgar.

FINANCIAL INSTRUMENTS

Financial instruments are defined as a contractual right or obligation to receive or deliver cash on another financial asset. The Corporation recognizes financial instruments based on their classification. Depending on the financial instrument's classification, changes in subsequent measurements are recognized in net loss or other comprehensive loss.

A description of the financial instruments, their fair value and risk management is included in the Corporation's annual audited consolidated financial statements for the year ended December 31, 2019 filed on SEDAR www.sedar.com and included in the registration statement on Form 40-F filed on EDGAR at www.sec.gov/edgar.

(Signed) Frédéric Ors

Frédéric Ors
Chief Executive Officer

(Signed) Pierre Labbé

Pierre Labbé
Chief Financial Officer

May 14, 2020

FORM 52-109F2
CERTIFICATION OF INTERIM FILINGS
FULL CERTIFICATE

I, Frederic Ors, Chief Executive Officer of IMV Inc., certify the following:

1. **Review:** I have reviewed the interim financial report and interim MD&A (together, the “interim filings”) of IMV Inc. (the “issuer”) for the interim period ended March 31, 2020.
 2. **No misrepresentations:** Based on my knowledge, having exercised reasonable diligence, the interim filings do not contain any untrue statement of a material fact or omit to state a material fact required to be stated or that is necessary to make a statement not misleading in light of the circumstances under which it was made, with respect to the period covered by the interim filings.
 3. **Fair presentation:** Based on my knowledge, having exercised reasonable diligence, the interim financial report together with the other financial information included in the interim filings fairly present in all material respects the financial condition, financial performance and cash flows of the issuer, as of the date of and for the periods presented in the interim filings.
 4. **Responsibility:** The issuer’s other certifying officer(s) and I are responsible for establishing and maintaining disclosure controls and procedures (DC&P) and internal control over financial reporting (ICFR), as those terms are defined in *National Instrument 52-109 Certification of Disclosure in Issuers’ Annual and Interim Filings*, for the issuer.
 5. **Design:** Subject to the limitations, if any, described in paragraphs 5.2 and 5.3, the issuer’s other certifying officer(s) and I have, as at the end of the period covered by the interim filings:
 - (a) designed DC&P, or caused it to be designed under our supervision, to provide reasonable assurance that:
 - (i) material information relating to the issuer is made known to us by others, particularly during the period in which the interim filings are being prepared; and
 - (ii) information required to be disclosed by the issuer in its annual filings, interim filings or other reports filed or submitted by it under securities legislation is recorded, processed, summarized and reported within the time periods specified in securities legislation; and
 - (b) designed ICFR, or caused it to be designed under our supervision, to provide reasonable assurance regarding the reliability of financial reporting and the preparation of financial statements for external purposes in accordance with the issuer’s GAAP.
 - 5.1 **Control framework:** The control framework the issuer’s other certifying officer(s) and I used to design the issuer’s ICFR is *Internal Control - Integrated Framework* issued by the Committee of Sponsoring Organizations of the Treadway Commission.
-

5.2 **ICFR - material weakness relating to design:** N/A

5.3 **Limitation on scope of design:** N/A

6. **Reporting changes in ICFR:** The issuer has disclosed in its interim MD&A any change in the issuer's ICFR that occurred during the period beginning on January 1, 2020 and ended on March 31, 2020 that has materially affected, or is reasonably likely to materially affect, the issuer's ICFR.

Date: May 15, 2020

(signed) Frederic Ors
Frederic Ors
Chief Executive Officer

FORM 52-109F2
CERTIFICATION OF INTERIM FILINGS
FULL CERTIFICATE

I, Pierre Labbé, Chief Financial Officer of IMV Inc., certify the following:

1. **Review:** I have reviewed the interim financial report and interim MD&A (together, the “interim filings”) of IMV Inc. (the “issuer”) for the interim period ended March 31, 2020.
 2. **No misrepresentations:** Based on my knowledge, having exercised reasonable diligence, the interim filings do not contain any untrue statement of a material fact or omit to state a material fact required to be stated or that is necessary to make a statement not misleading in light of the circumstances under which it was made, with respect to the period covered by the interim filings.
 3. **Fair presentation:** Based on my knowledge, having exercised reasonable diligence, the interim financial report together with the other financial information included in the interim filings fairly present in all material respects the financial condition, financial performance and cash flows of the issuer, as of the date of and for the periods presented in the interim filings.
 4. **Responsibility:** The issuer’s other certifying officer(s) and I are responsible for establishing and maintaining disclosure controls and procedures (DC&P) and internal control over financial reporting (ICFR), as those terms are defined in *National Instrument 52-109 Certification of Disclosure in Issuers’ Annual and Interim Filings*, for the issuer.
 5. **Design:** Subject to the limitations, if any, described in paragraphs 5.2 and 5.3, the issuer’s other certifying officer(s) and I have, as at the end of the period covered by the interim filings:
 - (a) designed DC&P, or caused it to be designed under our supervision, to provide reasonable assurance that:
 - (i) material information relating to the issuer is made known to us by others, particularly during the period in which the interim filings are being prepared; and
 - (ii) information required to be disclosed by the issuer in its annual filings, interim filings or other reports filed or submitted by it under securities legislation is recorded, processed, summarized and reported within the time periods specified in securities legislation; and
 - (b) designed ICFR, or caused it to be designed under our supervision, to provide reasonable assurance regarding the reliability of financial reporting and the preparation of financial statements for external purposes in accordance with the issuer’s GAAP.
 - 5.1 **Control framework:** The control framework the issuer’s other certifying officer(s) and I used to design the issuer’s ICFR is *Internal Control - Integrated Framework* issued by the Committee of Sponsoring Organizations of the Treadway Commission.
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5.2 **ICFR - material weakness relating to design:** N/A

5.3 **Limitation on scope of design:** N/A

6. **Reporting changes in ICFR:** The issuer has disclosed in its interim MD&A any change in the issuer's ICFR that occurred during the period beginning on January 1, 2020 and ended on March 31, 2020 that has materially affected, or is reasonably likely to materially affect, the issuer's ICFR.

Date: May 15, 2020

(signed) Pierre Labbé
Pierre Labbé
Chief Financial Officer
