



Condensed Interim Consolidated Financial Statements
For the three months ended March 31, 2025 and 2024

Notice of No Auditor Review of Unaudited Condensed Interim Consolidated Financial Statements

The accompanying unaudited condensed interim consolidated financial statements of Resverlogix Corp. (the “Company”) as at March 31, 2025 and for the period then ended have been prepared by and are the responsibility of the Company's management. The Company's Audit Committee and Board of Directors have reviewed and approved these unaudited condensed interim consolidated financial statements. In accordance with National Instrument 51 – 102, the Company discloses that its auditors have not reviewed the accompanying unaudited condensed interim consolidated financial statements for the periods ended March 31, 2025 and 2024.

Condensed Interim Consolidated Statements of Financial Position

As at:

(unaudited)

| <i>In thousands of US dollars</i> | Notes | March 31, 2025 | December 31, 2024 |
|---|-------|-------------------|----------------------|
| Assets | | | |
| Current assets: | | | |
| Cash | | \$ 153 | \$ 99 |
| Prepaid expenses and deposits | | 138 | 162 |
| Investment tax credit receivable | | 60 | 48 |
| Other assets | | 74 | 40 |
| Clinical supplies | | 936 | 996 |
| Due from Zenith Capital Corp. | | 391 | 193 |
| Total current assets | | 1,752 | 1,538 |
| Non-current assets: | | | |
| Intangible assets | | 2,615 | 2,487 |
| Clinical supplies | | 3,319 | 3,319 |
| Total non-current assets | | 5,934 | 5,806 |
| Total assets | | \$ 7,686 | \$ 7,344 |
| Liabilities | | | |
| Current liabilities: | | | |
| Trade and other payables | | \$ 16,294 | \$ 16,002 |
| Accrued interest | 5 | 2,393 | 1,781 |
| Promissory notes payable to Zenith Capital Corp. | 5 | 8,644 | 7,538 |
| Other promissory notes | | 759 | 759 |
| Lease liability | | 42 | 42 |
| Warrant liability | 7 (e) | 118 | 189 |
| Total current liabilities | | 28,250 | 26,311 |
| Non-current liabilities: | | | |
| Debt | | 8,131 | 8,131 |
| Other long-term liability | | 866 | 837 |
| Royalty preferred shares | 6 | 57,100 | 53,800 |
| Total liabilities | | 94,347 | 89,079 |
| Shareholders' deficiency: | | | |
| Share capital | 7 (a) | 334,716 | 334,617 |
| Contributed surplus | | 53,686 | 53,710 |
| Deficit | | (475,063) | (470,062) |
| Total shareholders' deficiency | | (86,661) | (81,735) |
| Total liabilities and shareholders' deficiency | | \$ 7,686 | \$ 7,344 |

Going concern (note 3)

Commitments and contingencies (note 9)

Signed on behalf of the Board:

Signed: "Kenneth Zuerblis" Director

Signed: "Kelly McNeill" Director

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

Condensed Interim Consolidated Statements of Comprehensive Loss (Income)
For the three months ended March 31
(unaudited)

| <i>In thousands of US dollars</i> | Notes | 2025 | 2024 |
|---|-------|-----------------|-------------------|
| Expenses: | | | |
| Research and development, net of recoveries | 8 | \$ 699 | \$ 681 |
| Investment tax credits | | (12) | (12) |
| Net research and development | | 687 | 669 |
| General and administrative, net of recoveries | 8 | 435 | 441 |
| | | 1,122 | 1,110 |
| Finance costs (income): | | | |
| Gain on change in fair value of warrant liability | 7 (e) | (71) | (121) |
| Loss (gain) on change in fair value of royalty preferred shares | 6 | 3,300 | (2,800) |
| Gain on change in fair value of derivative liability | | - | (163) |
| Interest and accretion | | 645 | 644 |
| Foreign exchange loss (gain) | | 2 | (13) |
| Net finance costs (income) | | 3,876 | (2,453) |
| Loss (income) before income taxes | | 4,998 | (1,343) |
| Income taxes | | 3 | 3 |
| Net and total comprehensive loss (income) | | \$ 5,001 | \$ (1,340) |
| Net loss (income) per share (note 7 (f)) | | | |
| Basic and diluted | | \$ 0.02 | \$ (0.00) |

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

Condensed Interim Consolidated Statements of Changes in Shareholders' Deficiency
For the three months ended March 31
(unaudited)

| <i>In thousands of US dollars</i> | Share Capital | Contributed Surplus | Deficit | Total Shareholders' Deficiency |
|---|--------------------------|--------------------------------|----------------|---|
| Balance, December 31, 2023 | \$ 333,716 | \$ 54,237 | \$ (462,427) | \$ (74,474) |
| Common shares issued in connection with long term incentive plan | 149 | (149) | - | - |
| Common shares issued in connection with deferred share unit plan | 141 | (141) | - | - |
| Share-based payment transactions | - | 42 | - | 42 |
| Net and total comprehensive income | - | - | 1,340 | 1,340 |
| Balance, March 31, 2024 | \$ 334,006 | \$ 53,989 | \$ (461,087) | \$ (73,092) |
| Balance, December 31, 2024 | \$ 334,617 | \$ 53,710 | \$ (470,062) | \$ (81,735) |
| Common shares issued in connection with long term incentive plan | 99 | (99) | - | - |
| Share-based payment transactions | - | 75 | - | 75 |
| Net and total comprehensive loss | - | - | (5,001) | (5,001) |
| Balance, March 31, 2025 | \$ 334,716 | \$ 53,686 | \$ (475,063) | \$ (86,661) |

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

Condensed Interim Consolidated Statements of Cash Flows
For the three months ended March 31
(unaudited)

| <i>In thousands of US dollars</i> | 2025 | 2024 |
|---|---------------|----------------|
| Cash provided by (used in): | | |
| Cash flows (used in) provided by operating activities: | | |
| Net (loss) income | \$ (5,001) | \$ 1,340 |
| Items not involving cash: | | |
| Equity-settled share-based payment transactions | 75 | 42 |
| Depreciation and amortization | 104 | 84 |
| Gain on change in fair value of warrant liability | (71) | (121) |
| Loss on change in fair value of royalty preferred shares | 3,300 | (2,800) |
| Gain on change in fair value of derivative liability | - | (163) |
| Unrealized foreign exchange | - | (5) |
| Interest and accretion | 645 | 644 |
| Net current income taxes | 3 | 3 |
| Changes in non-cash working capital: | | |
| Prepaid expenses and deposits | 24 | (720) |
| Investment tax credit receivable | (12) | (11) |
| Other assets | (34) | (48) |
| Clinical supplies | 60 | - |
| Due from Zenith Capital Corp. | (198) | (224) |
| Trade and other payables | 176 | 167 |
| | (929) | (1,812) |
| Interest received | - | 1 |
| Net cash used in operating activities | (929) | (1,811) |
| Cash flows provided by financing activities: | | |
| Proceeds from promissory notes payable | 1,106 | 2,000 |
| Net cash provided by financing activities | 1,106 | 2,000 |
| Cash flows used in investing activities: | | |
| Intangible asset additions | (232) | (182) |
| Changes in non-cash investing working capital | 109 | 44 |
| Net cash used in investing activities | (123) | (138) |
| Effect of foreign currency translation on cash | - | (1) |
| Increase in cash | 54 | 50 |
| Cash, beginning of period | 99 | 5 |
| Cash, end of period | \$ 153 | \$ 55 |

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

Notes to the Condensed Interim Consolidated Financial Statements

For the three months ended March 31, 2025 and 2024

(unaudited)

(Tabular amounts in thousands of US dollars, except for number of shares)

1. General information

Resverlogix Corp. (the “Company”) is a company domiciled in Canada. The annual consolidated financial statements comprise the Company and its wholly-owned subsidiary Resverlogix Inc. (together referred to as “Resverlogix” or the “Group”). Resverlogix Corp. is incorporated under the laws of Alberta. Resverlogix Inc. is incorporated under the laws of Delaware. The Company’s head office is located at Suite 300, 4820 Richard Road S.W., Calgary, Alberta, T3E 6L1. The registered and records office is located at Suite 600, 815 - 8th Avenue S.W., Calgary, Alberta, T2P 3P2.

Resverlogix is developing apabetalone (RVX-208), a first-in-class, small molecule that is a selective BET (bromodomain and extra-terminal) inhibitor. BET bromodomain inhibition is an epigenetic mechanism that can regulate disease-causing genes. Apabetalone is a BET inhibitor selective for the second bromodomain (“BD2”) within the BET proteins. This selective inhibition of apabetalone on BD2 produces a specific set of biological effects with potentially important benefits for patients with chronic disease including cardiovascular disease (“CVD”) and associated comorbidities, and post-COVID-19 conditions. Apabetalone is the only selective BET bromodomain inhibitor in human clinical trials. Apabetalone was studied in a Phase 3 trial, BETonMACE, in 13 countries worldwide, in high-risk CVD patients with type 2 DM and low high-density lipoprotein (“HDL”). The Company’s Phase 3 trial, BETonMACE, did not meet its primary endpoint but generated encouraging positive results in key secondary endpoints and the Company intends to continue the development of apabetalone when the requisite funding can be secured. Based on the results of the BETonMACE study, the U.S. Food and Drug Administration (“FDA”) granted Breakthrough Therapy Designation (“BTD”) for apabetalone in combination with top standard of care, including high-intensity statins, for the secondary prevention of major adverse cardiac events (“MACE”) in patients with type 2 DM and recent acute coronary syndrome (“ACS”). The achievement of BTD has the potential to expedite apabetalone’s clinical development program through more intensive FDA guidance. The Company is considered to be in the development stage, as most of its efforts have been devoted to research and development and it has not earned any revenue to date.

2. Basis of preparation

(a) Statement of compliance

These condensed interim consolidated financial statements have been prepared in accordance with IAS 34 – *Interim Financial Reporting*. These condensed interim consolidated financial statements were approved and authorized for issue by the Board of Directors on May 14, 2025.

(b) Basis of measurement

The condensed interim consolidated financial statements have been prepared on the historical cost basis except for liability classified warrants, liability classified royalty preferred shares and derivative liability, which are measured at fair value each reporting period.

(c) Functional and presentation currency

The functional currency of all entities within the Group is the US dollar, which is also the presentation currency. All financial information presented in dollars has been rounded to the nearest thousand except for per share amounts.

(d) Use of estimates and judgment

The preparation of the condensed interim consolidated financial statements in conformity with IFRS requires management to make judgments, estimates and assumptions that affect the application of accounting policies and the amounts reported in these condensed interim consolidated financial statements and notes. Accordingly, actual results may differ from estimated amounts as future confirming events occur. Significant estimates and judgment used in the preparation of the condensed interim consolidated financial statements remain unchanged from those described in the Group’s consolidated financial statements for the year ended December 31, 2024.

3. Going concern

The success of the Company is dependent on the continuation of its research and development activities, progressing the core technologies through clinical trials to commercialization or a strategic partnership, and its ability to obtain additional financing. It is not possible to predict the outcome of future research and development programs, the Company’s ability to fund these programs in the future, or to secure a strategic partnership, or the commercialization of products by the Company. To date, the Company has not generated any product revenue.

The consolidated financial statements have been prepared pursuant to International Financial Reporting Standards (“IFRS”) applicable to a going concern, which contemplates the realization of assets and settlement of liabilities in the normal course of

Notes to the Condensed Interim Consolidated Financial Statements

For the three months ended March 31, 2025 and 2024

(unaudited)

(Tabular amounts in thousands of US dollars, except for number of shares)

3. Going concern (continued)

business as they come due. The Company has incurred significant losses to date, and with no assumption of revenues, is dependent on its ability to raise additional financial capital by continuing to demonstrate the successful progression of its research and development activities if it is to remain as a going concern.

As at March 31, 2025, the Company had \$0.2 million of cash. The Company needs to raise additional capital to fund research, development and corporate activities over the next year or it may be forced to cease operations. As at March 31, 2025, the Company was committed to pay \$16.3 million of current trade and other payables (including \$11.7 million owing to EVERSANA Life Science Services, LLC ("EVERSANA")), \$8.6 million to Zenith Capital Corp. (a related party) ("Zenith") (due four months following demand), \$0.8 million of other unsecured promissory notes (due upon demand or four months following demand, respectively), up to \$0.2 million for research and development commitments, and \$0.2 million of operating lease expense over the next twelve months. The Company also has other commitments as outlined in Note 9. In addition, expenditures over the next twelve months under a cancellable agreement with a contract research organization in respect of planned clinical development are estimated to total approximately \$2-3 million.

The Company's cash as at March 31, 2025 is not sufficient to fund the Company's contractual commitments or the Company's planned business operations over the next year. During the three months ended March 31, 2025, Zenith advanced the Company \$1.1 million; there is no assurance that Zenith will advance further amounts to the Company. The Company will have to raise additional capital to fund its contractual commitments and its planned business operations. The Company continues to pursue and/or examine several sources of additional capital including co-development, licensing, rights or other partnering arrangements, procurement arrangements, private placements and/or public offerings (equity and/or debt). However, there is no assurance that any of these measures will be successful. The Company will also require additional capital to fund research, development and corporate activities beyond the next year. The Company will continue to explore alternatives to generate additional cash including raising additional equity and/or debt and/or partnering; however, there is no assurance that these initiatives will be successful.

These conditions result in a material uncertainty which may cast significant doubt on the Company's ability to continue as a going concern. If the Company is not able to raise capital, the Company may be forced to cease operations. These consolidated financial statements do not include necessary adjustments to reflect the recoverability and classification of recorded assets and liabilities and related expenses that might be necessary should the Company be unable to continue as a going concern and therefore be required to realize its assets and liquidate its liabilities and commitments in other than the normal course of business and such adjustments could be material.

4. Material accounting policies

The condensed interim consolidated financial statements should be read in conjunction with the Company's annual consolidated financial statements for the year ended December 31, 2024 prepared in accordance with IFRS applicable to those annual consolidated financial statements. The same accounting policies, presentation and methods of computation have been followed in these condensed interim consolidated financial statements as were applied in the Company's consolidated financial statements for the year ended December 31, 2024.

5. Promissory notes payable to Zenith Capital Corp.

The Company and Zenith have several directors and officers in common, and thus are considered related parties. The Company provides management and administrative services to Zenith pursuant to a Management Services Agreement dated June 3, 2013 between the Company and Zenith. The purpose of the agreement is to enable the Company to achieve greater utilization of its resources. As consideration for the services, Zenith pays the Company a service fee, consisting of salary and other compensation costs attributable to the services and reimbursable expenses incurred by Resverlogix in connection with the services.

During the three months ended March 31, 2025, the Company provided an aggregate of \$0.19 million (2024 – \$0.21 million) of services and reimbursable expenses, comprised of \$0.13 million (2024 – \$0.16 million) for management and administrative services, and \$0.08 million (2024 – \$0.07 million) of reimbursable expenses, less \$0.02 million (2024 – \$0.02 million) for services provided to Resverlogix by Zenith. The reimbursable expenses include proportionate share of rental payments and operating costs (for a laboratory and office that Resverlogix shares with Zenith) pursuant to a sublease that Resverlogix has in place with Zenith. Zenith owes the Company \$0.4 million (December 31, 2024 – \$0.2 million); this balance is unsecured, payable on demand and non-interest bearing.

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For the three months ended March 31, 2025 and 2024

(unaudited)

(Tabular amounts in thousands of US dollars, except for number of shares)

5. Promissory notes payable to Zenith Capital Corp. (continued)

During the three months ended March 31, 2025, Zenith advanced the Company \$1.1 million (2024 – \$2.0 million). The Company has issued promissory notes to Zenith totaling \$8.6 million as at March 31, 2025 (December 31, 2024 – \$7.5 million); the promissory notes bear interest at 12% per annum, are payable within four months of demand and are unsecured. During the three months ended March 31, 2025, interest on the Promissory notes payable to Zenith Capital Corp. totaled \$0.2 million (2024 – \$0.1 million) and is included in Accrued interest in the Consolidated Statements of Financial Position.

6. Royalty preferred shares

(i) Authorized:

Unlimited number of royalty preferred shares issuable in series with rights as determined by the Board of Directors at the time of issue.

(ii) Issued and outstanding:

| Preferred shares | Number of preferred shares | Amount |
|---|----------------------------|-----------|
| Balance, December 31, 2024 | 75,202,620 | \$ 53,800 |
| Revaluation of royalty preferred shares | - | 3,300 |
| Balance, March 31, 2025 | 75,202,620 | \$ 57,100 |

The holder of the royalty preferred shares is entitled to dividends in the amount of 6-12% of the Company's Net Revenue, as defined in the Company's articles. As at March 31, 2025, the Company had 75,202,620 royalty preferred shares outstanding, all of which were held by Zenith. Resverlogix and Zenith have several directors and officers in common, and thus are considered related parties. For fair value measurement purposes, the royalty preferred shares liability has been categorized within level 3 of the fair value measurement hierarchy. The estimated fair value of the royalty preferred shares is based on management's judgments, estimates and assumptions which include significant unobservable inputs including the timing and amounts of the Company's discounted future net cash flows. The estimate incorporates the following assumptions: an average cumulative probability rate of generating forecasted future cash flows of 41% as at March 31, 2025 (December 31, 2024 – 41%) reflecting in each case, among other factors, the Company's clinical results, in particular the results of BETonMACE, and communication with the U.S. Food and Drug Administration ("FDA") and other regulatory bodies; a discount rate of 24.3% as at March 31, 2025 (December 31, 2024 – 24.3%); projected commencement of revenue beginning between mid-2028 and early-2029 (based on projected clinical development paths across various jurisdictions, which is based substantially on securing the requisite funding from a partnership or other source(s) of capital by mid-2025) as at March 31, 2025 (December 31, 2024 – between mid-2028 and late-2028); and projected apabetalone market share percentages and projected product pricing. The estimated fair value of royalty preferred shares in the current period was affected by the passage of time (to future cash flows based on the estimated timing and commencement of revenue), offset by the commencement of revenue estimation update.

The estimated fair value of the royalty preferred shares is subject to significant volatility. Small changes in the aforementioned assumptions may have a significant impact on the estimated fair value of the royalty preferred shares. For instance, holding all other assumptions constant: a 1% increase in the discount rate would result in a \$4.0 million decrease in the estimated fair value of the royalty preferred shares; assuming commencement of revenue one year later would result in a \$16.7 million decrease in the estimated fair value of the royalty preferred shares; a 0.5% increase in the growth rate of the patient population would result in a \$4.3 million increase in the estimated fair value of the royalty preferred shares; and a 1% increase in the probability rate of generating forecasted future cash flows would result in a \$1.6 million increase in the estimated fair value of the royalty preferred shares.

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For the three months ended March 31, 2025 and 2024

(unaudited)

(Tabular amounts in thousands of US dollars, except for number of shares)

7. Shareholders' deficiency

(a) Common shares

(i) Authorized:

Unlimited number of common shares

(ii) Issued and outstanding:

| Common shares | Number of shares | Amount |
|--|------------------|------------|
| Balance, December 31, 2024 | 281,536,994 | \$ 334,617 |
| Issued in connection with long term incentive plan | 1,960,000 | 99 |
| Balance, March 31, 2025 | 283,496,994 | \$ 334,716 |

(b) Stock options

The Company's amended stock option plan has been approved as a rolling 10% plan that allows for reservation of a number of common shares under the plan equal to 10% of the Company's issued and outstanding common shares on an undiluted basis. Additionally, the plan is a reloading plan, which allows for the number of common shares reserved for issuance related to the options under the plan to automatically become eligible to be reallocated pursuant to stock option-based grants upon option expiry, cancellation or exercise. The Company may grant options to its directors, officers, employees and consultants. The majority of options fully vest over one to three years and have a five-year term. The options are settled by way of the issuance of equity instruments of the Company ("equity-settled").

| | Number of options | Weighted average exercise price (CAD) |
|---|-------------------|---------------------------------------|
| Outstanding, December 31, 2024 and March 31, 2025 | 2,630,000 | \$ 0.24 |

The fair value of each option granted is estimated as of the grant date using the Black-Scholes option pricing model. The following weighted average assumptions were used in arriving at the weighted average fair values of \$0.04 per option associated with stock options granted during the three months ended March 31, 2024:

| | 2024 |
|---------------------------|-----------|
| Risk-free interest rate | 3.7% |
| Expected life | 4.3 years |
| Expected volatility | 90% |
| Share price at grant date | CAD\$0.07 |
| Expected dividends | Nil |

The following table summarizes information about the options outstanding and exercisable at March 31, 2025.

| Range of Exercise Prices (CAD) | Number Outstanding | Weighted Average Remaining Life (years) | Weighted Average Exercise Price (CAD) | Number Exercisable |
|--------------------------------|--------------------|---|---------------------------------------|--------------------|
| \$0.07 - \$0.18 | 2,040,000 | 3.17 | \$ 0.10 | 1,965,000 |
| \$0.54 - \$0.72 | 390,000 | 1.82 | 0.65 | 390,000 |
| \$0.79 - \$0.91 | 200,000 | 0.91 | 0.88 | 200,000 |
| | 2,630,000 | 2.80 | \$ 0.24 | 2,555,000 |

The number of options exercisable at March 31, 2025 was 2,555,000 (2024 - 2,655,000) with a weighted average exercise price of CAD\$0.25 (2024 - CAD\$0.34).

Notes to the Condensed Interim Consolidated Financial Statements

For the three months ended March 31, 2025 and 2024

(unaudited)

(Tabular amounts in thousands of US dollars, except for number of shares)

7. Shareholders' deficiency (continued)

(c) Restricted stock units

The Company's long term incentive plan allows for the reservation of a number of common shares not to exceed 10% of the Company's issued and outstanding common shares on an undiluted basis less the number of common shares reserved under the Company's amended stock option plan. The Company may grant restricted stock units ("RSUs") to directors, officers, employees, and consultants. RSUs are settled on exercise through the issuance of common shares.

During the three months ended March 31, 2025, 1,400,000 RSUs were granted (2024 – 1,230,000 RSUs were granted). The RSUs vest over a period of zero to six months. The Company estimates the fair value of RSUs based on the market price of the underlying stock on the date of grant.

| | Number of restricted stock units | Weighted average grant date fair value (USD) |
|--------------------------------|-------------------------------------|---|
| Outstanding, December 31, 2024 | 19,790,075 | \$ 0.58 |
| Granted | 1,400,000 | 0.03 |
| Exercised | (1,960,000) | 0.05 |
| Outstanding, March 31, 2025 | 19,230,075 | \$ 0.60 |

At March 31, 2025, 18,655,073 RSUs were exercisable (2024 – 20,209,161).

(d) Deferred share units

The Company's deferred share unit plan limits the maximum number of Common Shares issuable pursuant to outstanding deferred share units ("DSUs") at any time to 5% of the aggregate number of issued and outstanding Common Shares, provided that the combined maximum number of Common Shares issuable by the Company pursuant to outstanding DSUs and all of its other security-based compensation arrangements may not exceed 10% of the Common Shares outstanding from time to time. The Company may grant DSUs to directors. DSUs are settled on exercise through the issuance of common shares.

During the three months ended March 31, 2025, there were no DSUs granted to directors (2024 – no DSUs were granted). The DSUs fully vest at grant date. The Company estimates the fair value of DSUs based on the market price of the underlying stock on the date of grant.

| | Number of deferred share units | Weighted average grant date fair value (USD) |
|---|-----------------------------------|---|
| Outstanding and exercisable, December 31, 2024 and March 31, 2025 | 3,431,420 | \$ 0.20 |

(e) Warrant liability

The following table summarizes the changes in liability-classified common share purchase warrants outstanding.

| | Number of warrants | Weighted average exercise price (CAD) | Liability amount |
|----------------------------------|-----------------------|--|---------------------|
| Outstanding, December 31, 2024 | 26,261,157 | \$ 0.21 | \$ 189 |
| Revaluation of warrant liability | - | - | (71) |
| Outstanding, March 31, 2025 | 26,261,157 | \$ 0.21 | \$ 118 |

The following table summarizes information about liability-classified warrants outstanding and exercisable at March 31, 2025.

| Exercise Price (CAD) | Number Outstanding and Exercisable | Weighted Average Remaining Life (years) | Weighted Average Exercise Price (CAD) |
|----------------------|---------------------------------------|--|--|
| \$0.20 | 25,961,157 | 1.61 | \$ 0.20 |
| \$0.93 | 300,000 | 0.12 | 0.93 |
| | 26,261,157 | 1.59 | \$ 0.21 |

Notes to the Condensed Interim Consolidated Financial Statements

For the three months ended March 31, 2025 and 2024

(unaudited)

(Tabular amounts in thousands of US dollars, except for number of shares)

7. Shareholders' deficiency (continued)

(e) Warrant liability (continued)

Under IFRS, the prescribed accounting treatment for warrants, with an exercise price denominated in a foreign currency, is to treat these warrants as a liability measured at fair value with subsequent changes in fair value each reporting period accounted for through profit or loss. The initial fair value of these warrants is determined using the Black Scholes option pricing model.

The Company's warrants are presented as a current liability on the consolidated statements of financial position. Each full warrant entitles the holder to purchase one common share of the Company. As these warrants are exercised, the fair value of the recorded warrant liability on the date of exercise is included in share capital along with the proceeds from the exercise. If these warrants expire, the related decrease in warrant liability is recognized in profit or loss, as part of the change in fair value of warrant liability.

The fair value of the warrants not publicly listed is determined using the Black Scholes option pricing model at initial issue date and at each reporting period, unless the warrants are listed, in which case the initial trading value is used.

The changes in fair value of the unlisted liability-classified warrants were based on several factors including changes in the market price of the Company's shares from CAD\$0.055 on December 31, 2024 to CAD\$0.04 on March 31, 2025, and from CAD\$0.07 on December 31, 2023 to CAD\$0.06 on March 31, 2024, as well as decreases in the remaining terms of the various series of warrants, and changes in estimated future volatility of our common shares which represents a level 3 input in the fair value hierarchy. The fair value of the warrants is subject to significant volatility. Gains and losses resulting from the revaluation of warrant liability are non-cash and do not impact the Company's cash flows.

There were no warrants issued during the three months ended March 31, 2025 (2024 – no warrants were issued).

(f) Per share amounts

The basic and diluted net (income) loss per share have been calculated based on the weighted average shares outstanding:

| | 2025 | 2024 |
|---|-------------|-------------|
| Weighted average common shares outstanding - basic and diluted | 282,774,105 | 273,668,456 |
| Effect of convertible debenture, warrants, stock options, RSUs and DSUs | - | 169,184,219 |
| Weighted average common shares outstanding - diluted | 282,774,105 | 442,852,675 |

The effect of any potential exercise of convertible debenture, warrants, stock options, restricted stock units, and deferred share units outstanding is excluded from the calculation of diluted loss per share in periods where the effect would be anti-dilutive.

8. Expenses by nature

Presentation of expenses is based on the function of each expense. The following details provide a breakdown of the components of the research and development and general and administrative expenses classified by nature.

| | 2025 | 2024 |
|--|--------|--------|
| Research and development expenses: | | |
| Operating expenses, net of recoveries | \$ 351 | \$ 344 |
| Personnel costs (short-term employee benefits) | 244 | 253 |
| Amortization and depreciation | 104 | 84 |
| Total research and development expenses | \$ 699 | \$ 681 |
| General and administrative expenses: | | |
| General expenses, net of recoveries | \$ 28 | \$ 54 |
| Personnel costs (short-term employee benefits) | 332 | 345 |
| Share-based payment transaction costs | 75 | 42 |
| Total general and administrative expenses | \$ 435 | \$ 441 |

Notes to the Condensed Interim Consolidated Financial Statements

For the three months ended March 31, 2025 and 2024

(unaudited)

(Tabular amounts in thousands of US dollars, except for number of shares)

9. Commitments and contingencies

As at March 31, 2025, the Group is committed to expenditures over the next twelve months of \$0.2 million (December 31, 2024 – \$1.6 million) under various research and development contracts. As at March 31, 2025, the Group is also party to a cancellable agreement with a contract research organization in respect of planned clinical development. Corresponding estimated aggregate expenditures over the next twelve months total approximately \$2-3 million (December 31, 2024 – \$2-3 million).

As at March 31, 2025, the Group is also party to a commercialization partnership with EVERSANA; the parties have mutually agreed to temporarily pause services and the Group was not obligated as at March 31, 2025 to incur pre-commercialization costs over the next twelve months. The parties may or may not resume services over the next twelve months.

The July 2015 License Agreement between Resverlogix and Hepalink was amended effective June 17, 2022, and further amended on June 27, 2024, such that Resverlogix agreed to pay up to CAD\$8.0 million of clinical development costs associated with apabetalone, including a global Phase 3 clinical trial (which Resverlogix intends to perform in any event), in China, Hong Kong, Taiwan and Macau, and if the costs incurred by Resverlogix after May 1, 2020 and up to June 30, 2025 total less than CAD\$8 million, then Resverlogix and Hepalink shall negotiate a mutually-agreeable timeframe regarding any difference, in principle by not later than June 30, 2026.

In July 2020, the Company entered into an agreement with a supplier to settle amounts owing by the Company, whereby the Company agreed to pay a reduced amount in three instalments of \$200,000, \$550,000 and \$550,000 on August 1, 2020, September 1, 2020 and October 1, 2020 respectively. The Company paid the August 1, 2020 instalment and has paid an additional \$825,000, but has not yet paid the remaining balance of \$275,000. Until the Company pays the remaining \$275,000, thereby satisfying its obligations pursuant to the agreement, it is possible that the supplier could assert that the Company is in default and could pursue any remedies that may be available to them.

The Company has not complied fully with the payment terms associated with certain amounts owing to certain vendors. Until the Company fully satisfies its obligations, it is possible that the vendors could assert that the Company is in default and could pursue any remedies available to them.

In 2021, the Company acquired certain intellectual property for: (a) \$400,000 paid in cash and (b) a \$600,000 milestone payment payable upon submission of a New Drug Application for apabetalone to the US Food and Drug Administration.