

Preliminary Final Report for the Period Ended 30 June 2015

Melbourne, Australia, 24 August 2015 - Antisense Therapeutics Limited ("ASX.ANP" or "the Company") is pleased to release its Preliminary Final Report for the year ended 30 June 2015.

During the year Antisense Therapeutics has made substantial progress in the clinic with its advanced stage antisense product pipeline and with partnering for ongoing development and commercialization of its compounds. The Company ended the year with a strengthened balance sheet and cash at bank as at 30 June 2015 of \$6.8 million.

Key operational highlights include:

ATL1103 for Acromegaly, Diabetic Retinopathy and Nephropathy and Cancer

- Successful efficacy results from the Phase II trial for Acromegaly (ATL1103/COR-004);
- Executed license agreement with Cortendo AB for endocrinology applications;
- Progressing plans for entry into Phase III clinical development;
- Cortendo AB to fund ongoing development.

ANP received an upfront payment of US\$5m (AUD \$6.4m) with potential additional payments of up to US\$105m (contingent upon achieving specific clinical, regulatory and commercialization milestones) and royalty payments based on sales performance. Importantly, ANP retains worldwide rights for other ATL1103 indications and Australian and New Zealand commercialization rights for Acromegaly. ANP's licensing partner Cortendo AB is to fund associated future development, regulatory and drug manufacture costs for ATL1103/COR-004 for Acromegaly.

ATL1102 for Multiple Sclerosis (MS)

- Supportive guidance obtained from the US Food and Drug Administration (FDA) Pre-IND assessment of the development strategy for ATL1102, including plans for a Phase IIb study in MS patients;
- Engaged in a process to attract a development partner for further clinical development;
- Global agreement executed with myTomorrows (Amsterdam, The Netherlands) to implement an Early Access Program (EAP) for ATL1102 for the treatment of Multiple Sclerosis (MS).

myTomorrows will perform at their cost the EAP activities including relevant data collection and the seeking of the regulatory approvals. Subject regulatory approvals and support for the ATL1102 EAP program, ANP expects to provide ATL1102 to MS treatment centers in the EU at prices that are comparable to current medicines used to treat MS. Initially the focus will be on those major European countries where the drug would qualify for use.

Board changes

To support the continued development and growth of the Company, the Board has commenced the search for potential new Board members. The skills, experience and key attributes being sought are considered important for the next stage in the implementation of the drug development and commercialisation strategy of the Company. Further to this, and after 15 years on the Board, Dr Chris Belyea has indicated that he will not be standing for reelection at the planned 2015 Annual General Meeting. Dr. Belyea was a founding Director and CEO of the Company at the time of listing in 2001. His industry knowledge and scientific and commercial insights have been greatly valued.

Contact Information:

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Managing Director: Mark Diamond +61 (3) 9827 8999



Appendix 4E – Preliminary Final Report

(ASX Listing rule 4.2A)

Company Name: Antisense Therapeutics Limited (the 'Company')

ABN: 41 095 060 745

Reporting Period: Financial year ended 30 June 2015
Previous Reporting Period: Financial year ended 30 June 2014

Result for Announcement to the Market

The results of Antisense Therapeutics Limited for the year ended 30 June 2015 are as follows:

Revenues	up	4,622.12%	to	\$3,916,337
Profit after tax attributable to members	up	123.46%	to	\$706,918
Net profit for the period attributable to members	up	123.46%	to	\$706,918

Brief explanation of figures reported above

The profit for the Company after income tax for the reporting period was \$706,918 (2014: loss \$3,013,272) and before income tax the profit for the reporting period was \$706,918 (2014: loss \$3,013,272). This result has been achieved after fully expensing all research and development costs, in the current reporting period of \$1,675,820 (2014: \$2,146,463).

For further details relating to the current period's results, refer to the Operations Report contained within this document.

Dividends

No dividends have been paid or declared by the Company since the beginning of the current reporting period. No dividends were paid for the previous reporting period.

Net Tangible Assets per Share

	30 June 2015	30 June 2014
Net Tangible Assets	\$7,091,598	\$2,086,891
Shares (No.)	176,512,483	144,096,128
Net Tangible Assets per Share (Cents)	4.02	1.45

Earnings/(Loss) per Share

	30 June 2	015	30 June 2014
Basic earnings/(loss) per share		0.45	(2.09)
Diluted earnings/(loss) per share		0.45	(2.09)

Status of Audit of Accounts

This Appendix 4E is based on accounts which have been audited. The audit report is included within the financial report which accompanies this Appendix 4E.



Annual Financial Report For the Year Ended 30 June 2015

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Operations Report

Overview of Company's activities

Antisense Therapeutics Limited ("the Company" or "Antisense Therapeutics") continued its focus on advancing its antisense products under development. The following report on operations details the research and development activities undertaken by the Company in the period.

Antisense Therapeutics' Mission

Antisense Therapeutics' mission is to develop and commercialise novel antisense therapeutics in-licensed from Isis Pharmaceuticals Inc (Isis), world leaders in antisense drug discovery and development. The Company's Research and Development activities are focused on developing its pipeline of 2nd generation antisense drugs for diseases where there is a significant and acknowledged unmet medical need and where the antisense technology has the potential to provide compounds with clear competitive advantages over existing therapies or drugs in development for those diseases.

Antisense Technology

Antisense drugs are small (12-21 nucleotides) pieces of DNA or RNA that are chemically modified to engineer good drug properties. Conventional medicines typically bring about their desired therapeutic effect by binding to a target protein directly, to interfere with the action of the disease causing protein. Antisense drugs on the other hand, are rationally designed to bind to a specific messenger RNA sequence with extraordinary precision and thereby block or stop the production of the disease causing protein in the first instance.

The antisense drugs in our pipeline accessed via the Company's technology collaboration with Isis incorporate Isis second-generation chemistry. Second-generation drugs are composed of both RNA-like and DNA-like nucleotides, while first-generation drugs are entirely DNA-like. Because RNA hybridizes more tightly to RNA than to DNA, the second-generation drugs have a greater affinity for their RNA targets and, therefore, greater potency than their first generation antisense drugs. Second generation antisense drugs are more stable, allowing more convenient dosing regimens, better tolerated, and have broad disease application.

Projects Update

ATL1103 for Acromegaly, Diabetic Retinopathy and Nephropathy and Cancer

ATL1103 is a second generation antisense drug designed to block growth hormone receptor (GHr) expression thereby reducing levels of the hormone insulin-like growth factor-I (IGF-I) in the blood and is a potential treatment for diseases associated with excessive growth hormone action. By inhibiting GHr production, ATL1103 in turn reduces IGF-I levels in the blood (serum). There are a number of diseases that are associated with excess GH and IGF-I action. These diseases include acromegaly, an abnormal growth disorder of organs, face, hands and feet; diabetic retinopathy, a common disease of the eye and a major cause of blindness; diabetic nephropathy, a common disease of the kidney and major cause of kidney failure, and certain forms of cancer.

ATL1103 is in clinical development as a treatment for acromegaly. The therapeutic activity of ATL1103 has been demonstrated both in animal pharmacology studies, where ATL1103 has shown the successful suppression of serum IGF-I levels in both mice and primates, and in a Phase I clinical trial in healthy volunteers. Normalizing serum IGF-I levels is the therapeutic goal in the treatment of acromegaly and reducing the effects of IGF-I has a potential role in the treatment of diabetic retinopathy, nephropathy and certain forms of cancer.

Operations Report (continued...)

Following the successful Phase I trial, the Company initiated a Phase II clinical trial of ATL1103 in patients with acromegaly.

In September 2014 the Company reported successful efficacy results from the Phase II trial with the trial having met its primary efficacy endpoint by showing a statistically significant average reduction in sIGF-I levels of 26% from baseline (P<0.0001) at week 14 (one week past the last dose) at the 400mg per week dose tested. ATL1103 was assessed as generally well tolerated and the positive safety profile suggested that the drug may be tolerated at higher dose levels than 400mg per week.

In early December 2014 the Company advised that it was planning to undertake a small, higher dose (600mg/week) study in Australia in 4 acromegaly patients to support the use of a higher dose of ATL1103 in future Phase III trials for dose escalation in patients with more active disease. Later in the same month the Company advised that it had received Ethics Committee approval to conduct this higher dose trial of ATL1103.

Progress

On 5th March 2015 the Company advised that it had received the requisite approvals and acknowledgements to commence patient enrolment in its ATL1103 higher dose study. The design of this higher dose trial is an open-label study of the safety, tolerability, pharmacokinetics and efficacy [effect on serum insulin like growth factor I (sIGF-I)] of ATL1103 in adult patients with acromegaly dosed twice weekly with ATL1103 at 300mg for 13 weeks (600mg weekly) with two months of follow up.

On 6 March 2015 the Company announced that Dr Peter Trainer, Professor of Endocrinology, The Christie NHS Foundation Trust, UK, and Chief Investigator for the Company's ATL1103 Phase II study, would present on the ATL1103 project and the previously announced Phase II trial results at the world's largest endocrinology meeting, ENDO 2015.

On 15th May 2015 the Company announced that Cortendo AB a biopharmaceutical company focused on rare endocrine disorders and other rare diseases and Antisense Therapeutics Limited had entered into an exclusive license agreement that provides Cortendo with development and commercialization rights to Antisense Therapeutics' ATL1103 for endocrinology applications.

Under the terms of the agreement, Cortendo provided Antisense Therapeutics with an initial upfront payment of \$5 million (AUD \$6.4 million), consisting of \$3 million (AUD \$3.9 million) in cash and a \$2 million (AUD \$2.5 million) investment in Antisense Therapeutics equity. Additional payments, contingent upon achieving specific development and commercialization milestones, may total up to \$105 million (AUD \$131 million) over the lifetime of the agreement. There is also the potential for royalty payments based upon sales performance.

Cortendo will be responsible for the ongoing clinical development of ATL1103 in endocrinology applications and will fund the associated future development, regulatory and drug manufacture costs. Antisense Therapeutics will retain commercialization rights for ATL1103 in endocrinology applications in Australia and New Zealand, and will also retain worldwide rights for other ATL1103 indications, and may utilize new ATL1103 data generated by Cortendo in pursuing these other indications, subject to certain terms and conditions.

Patent News

On 7th July 2014 the Company announced that the Canadian Patent Office allowed patent application 2,517,101 entitled "A Modified Oligonucleotide for inhibition of Growth Hormone Receptor Expression" which covers the Company's growth hormone receptor (GHr) targeting drug ATL1103 and its use until February 2024.

On 16th April 2015 the Company announced that the European Patent Office had allowed European patent application 11194098.7 entitled "Modulation of growth hormone receptor expression and insulin-like growth



Operations Report (continued...)

factor expression" and provides protection in the major pharmaceutical markets in Europe to 2024, with potential for extension to 2029.

What is Acromegaly?

Acromegaly is a serious chronic life threatening disease triggered by excess secretion of growth hormone (GH) by benign pituitary tumours. Oversupply of GH over stimulates liver, fat and kidney cells, through their GH receptors, to produce excess levels of Insulin-Like Growth Factor-I (IGF-I) in the blood manifesting in abnormal growth of the face, hands and feet, and enlargement of body organs including liver, kidney and heart. The primary treatments for acromegaly are to surgically remove the pituitary gland and/or drug therapy to normalize GH and serum IGF-I levels. In North America and Europe there are approximately 85,000 diagnosed acromegaly patients with about half requiring drug therapy.

ATL1102 for Multiple Sclerosis (MS)

ATL1102 is a second generation antisense inhibitor of CD49d, the alpha subunit of VLA-4 (Very Late Antigen-4). In inflammation, white blood cells (leukocytes) move out of the bloodstream into the inflamed tissue, for example, the Central Nervous System (CNS) in MS, and the lung airways in asthma. In MS, the inhibition of VLA-4 prevents white blood cells from entering the CNS, thereby reducing the severity of the disease and slowing its progression. VLA-4 is a clinically validated target in the treatment of MS. Antisense inhibition of VLA-4 has demonstrated positive effects in a number of animal models of inflammatory disease including MS. ATL1102 was shown to be highly effective in reducing MS lesions in a Phase IIa clinical trial in MS patients.

In September 2014 the Company reported the publication of previously generated Phase IIa clinical trial data on ATL1102 in the medical journal *Neurology*. The article titled "CD49d antisense drug ATL1102 reduces disease activity in patients with relapsing-remitting MS", was included in the print edition Volume 83, November 11, 2014.

In October 2014 the Company reported that the US Food and Drug Administration (FDA) had responded affirmatively to the Company's plan to submit a U.S. Investigational New Drug (IND) application for initiation of longer term Phase IIb human trials of ATL1102 for the treatment of Multiple Sclerosis (MS) and that supportive guidance had been obtained from the agency's Pre-IND assessment of the development strategy for ATL1102, including plans for a Phase IIb study in MS patients.

In December 2014 the Company reported that it was investigating provision of ATL1102 under an Early Access Program (EAP) on compassionate use or on a named patient basis in markets where the drug would qualify for use on these grounds including those where the Company can charge for drug access resulting in a possible early income stream and that Antisense Therapeutics was in discussions with an experienced European based group to set up and run the program in Europe for the Company.

Progress

On 27th May 2015 the Company announced that it had signed a global agreement with innovative expanded access provider myTomorrows (Amsterdam, The Netherlands) to implement an Early Access Program (EAP) for ATL1102 for the treatment of Multiple Sclerosis (MS). Subject to myTomorrows receiving the requisite regulatory approvals and support for the ATL1102 EAP program, ANP expects to provide ATL1102 to MS treatment centres in the EU at prices that are comparable to current medicines used to treat MS. Initially the focus will be on those major European countries where the drug would qualify for use. Under the EAP agreement, myTomorrows will perform at their cost the EAP activities including relevant data collection and the seeking of the EAP approvals. myTomorrows are to receive a share of EAP related revenue less the cost of drug and associated pass through costs including those to Isis Pharmaceuticals from whom ANP in-licensed ATL1102.



Operations Report (continued...)

ANP is working to firm up suitable ATL1102 drug product supply and the initial quantities for use in the program while its partner, myTomorrows, is preparing documentation for physician education and the EAP approvals process.

Patent News

On 16th September 2014 the Company announced that the European Patent Office had allowed European patent application 09798248.2, entitled "Methods for Treating Multiple Sclerosis using Antisense Oligonucleotides" which extends coverage of the ATL1102 compound for the treatment of relapsing-remitting multiple sclerosis (RRMS) patients until 2029 with potential for up to 5 year extension to 2034. With this allowance the granting of the European patent is a formality and will take place in the coming months.

What is Multiple Sclerosis?

Multiple Sclerosis (MS) is a life-long, chronic disease that progressively destroys the central nervous system (CNS). It affects approximately 400,000 people in North America and more than 1 million worldwide and the current market for MS drugs is estimated at more than USD\$12 billion. It is a disease that affects more women than men, with onset typically occurring between 20 and 40 years of age. Symptoms of MS may include vision problems, loss of balance, numbness, difficulty walking and paralysis. In Australia MS affects over 15,000 people and worldwide MS may affect more than one million people.

ATL1102 for Stem Cell Mobilization

Stem cell transplantation is a medical procedure used to improve clinical outcomes for patients undergoing chemotherapy to treat cancer. The Company identified a potential application for ATL1102 as a stem cell mobilization agent for use in combination with G-CSF (the main agent used for hematopoietic stem cell mobilization) in stem cell transplantation. In July the Company announced the results of the proof of concept trial which showed that use of ATL1102 in combination with G-CSF did not appear to increase the release of CD34+ stem cells beyond that achieved with G-CSF alone and consequently was not planning to move forward with the clinical development of ATL1102 in the SCM indication.

ATL1102 for Asthma

The Company has previously reported encouraging results achieved in an animal model of asthma with the inhaled form of an antisense compound targeting the VLA-4 molecule. Experimental studies showed that the delivery of an antisense drug against VLA-4 via inhalation to the lung significantly suppressed the key asthma indicators in allergen sensitized mice at very low inhaled doses, pointing to the potential application of ATL1102 as an inhaled treatment for asthma. The Company has conducted successful animal studies using inhaled ATL1102. Further development for the inhaled asthma application of ATL1102 would be undertaken with a partner.

ATL1101 for Prostate Cancer

ATL1101 is an antisense inhibitor of insulin like growth factor 1 receptor (IGF-Ir). IGF-Ir is one of the best known of a family of cell signalling molecules that are referred to as "anti-apoptotic". These molecules prolong cell survival by inhibiting programmed cell death (apoptosis). Inhibition of cell survival molecules like IGF-Ir can render tumour cells more susceptible to cell death with cytotoxic (cell death inducing) drugs. Similar "chemosensitiser" therapeutic approaches targeting the IGF-Ir are under investigation in several large pharmaceutical companies, lending support to ATL's antisense-based strategy against the same target. In animal studies ATL1101 demonstrated its effectiveness in suppressing human prostate cancer tumour growth in mouse models of human prostate cancer. ATL has previously undertaken certain toxicology studies on ATL1101 that would potentially position the drug to move into a clinical study in patients with prostate cancer. Further clinical development of ATL1101 would be anticipated to occur with a partner.



What is Prostate Cancer?

Prostate cancer is the second most frequently diagnosed cancer in men after skin cancer. Metastatic disease invariably progresses to hormone refractory or castrate-resistant prostate cancer (CRPC) if given enough time. Prostate tumours are initially androgen (male sex hormone) dependent, and can be treated with androgen ablation therapy (the term "castration" can be used to describe removal of the source of androgen), however once the disease progresses to its most dangerous and aggressive form, CRPC, treatment options are limited and prognosis is poor. Treatment options depend on disease severity and include radiation and chemotherapy, which are designed to induce programmed cell death (apoptosis) of tumour cells. There is a pressing need for the development of new treatment options for CRPC.

R&D Tax Incentive

During the year the Company received from the ATO a payment of \$1,139,739 in relation to R&D expenditure incurred in the 30 June 2014 financial year.

Capital Raising

Through October and November 2014, the Company raised \$2 million through a placement to institutional investors and a share purchase plan.

As referred to in the Projects Update section of the report, the Company received \$2.5 million through a share issue to license partner Cortendo. These shares are held in escrow for up to 2 years.

For further details, please refer to Note 16 Contributed Equity.

Financial Position

At 30 June 2015, the Company had cash reserves of \$6,829,605 (2014: \$1,334,513).

Events after Balance Sheet Date

No matters or circumstances have arisen since the end of the reporting period, not otherwise disclosed in this report, which significantly affected, or may significantly affect, the operations of the Company, the result of those operations, or the state of affairs of the Company in subsequent financial periods.

Auditor's Independence Declaration

A copy of the Auditor's Independence Declaration as required under section 307C of the *Corporations Act 2001* is set out on the page 33.



Intellectual Property Report

Antisense Therapeutics currently has 8 patent families with 63 patents registered and 12 patent applications pending covering its three antisense drugs ATL1101, ATL1102, and ATL1103 and their applications. Antisense Therapeutics has also licensed from Isis Pharmaceuticals, 19 Isis proprietary patents and applications directed to the antisense drug platform together with rights to 11 other Isis manufacturing patent families.

Since reporting on the status of the Company's intellectual property portfolio in the 2014 Annual Report the Company has successfully expanded its patent portfolio as follows:

- 2 US patents and a Canadian patent and a key European patent have been issued and/or registered;
 - O US 8,637,484, covering 48 other antisense to human GHr that reduce GHr has been registered and Canadian patent 2,517101 covering ATL1103 has been granted;
 - European application 09798248.2 covering ATL1102 in the treatment of relapsing forms of MS has been granted and registered in 10 European countries; and
 - US 9,084,770 covering the use of the ATL1101 compound and other antisense to IGF-IR for enhancing the sensitivity of IGF-IR positive tumors or cancer cells to a taxane has been issued;
- A US continuation application 14/731203 has been filed covering the use of ATL1101 and other antisense to IGF-IR to treat IGF-IR positive prostate tumors or cancers;
- A European divisional application of 09798248.2 has been filed covering the use of ATL1102 for the treatment of nonrelapsing forms of MS such as primary progressive and secondary progressive MS together with a corresponding Japanese Divisional.

The progress outlined above has added significant value to an already extensive intellectual property portfolio. Key patents have been granted for all of the compounds in Antisense Therapeutics' product pipeline that underpin Antisense Therapeutics commercialisation plans for its antisense drugs.

Country	Patent application or	Current Status	Expiry
	Patent No.		
ATL1103 Patent Portfolio	o **		
US	7,803,781	Patent Registered	2025*
US	8,299,039	Patent Registered	2024*
US	8,637,484	Registered	2024*
International	PCT/US2004/005896	National Phase applications	
Australia	2004217508	Patent Registered	2024*
Canada	2,517,101	Patent Registered	2024
Europe***	04715642.7	Under Examination	2024*
Europe***	11194098.7 Divisional of 04715642.7	Under Examination	2024*
Japan	2006-508878	Patent Registered	2024*
Japan	Divisional of 2006-508878	Under Examination	2024*
New Zealand	542595	Patent Registered	2024
USA	7,846,906	Patent Registered	2024*
USA	8,623,836	Patent Registered	2024*
USA	14/137852 Continuation of US/12/953105	Under Examination	2024*
International	PCT/AU2013/000095	National Phase Applications	
Australian	2013214698	Filed	2032
Canada	2863499	Filed	2032



Intellectual Property Report (continued...)

Europe	13743020.3	Filed	2032
Japan	2014-555044	Filed	2032
New Zealand	629004	Under Examination	2032
USA	14/376390	Under Examination	2032
International	PCT/AU2014/000613	International Phase	2033
ATL1102 Patent Portfolio	**		
USA	US 5968 826	Patent Registered	2018 **
USA	US 6258 790	Patent Registered	2018*/**
International	PCT/US99/18796	National Phase applications	
Australia	AU 759938	Patent Registered	2019 *
Canada	2,345,209	Patent Registered	2019
Japan	2000-574727	Patent Registered	2019*
Japan	2006-000258	Patent Registered	2019*
Europe	EP1123414	Regional Phase - granted	
Denmark	DK/EP1123414	Patent Registered	2019*
Finland	EP(FI)1123414	Patent Registered	2019*
France	EP(FR)1123414	Patent Registered	2019*
Germany	DE69934998.2-08	Patent Registered	2019*
Italy	IT40051BE2007	Patent Registered	2019*
Spain	ES2279632	Patent Registered	2019*
Sweden	SE99942290.0	Patent Registered	2019*
United Kingdom	EP(UK)1123414	Patent Registered	2019*
ATL1102 MS Patent Portf	olio **		
International	PCT/US2009/003760	National Phase applications	
Australia	AU 2009271678	Patent Registered	2029*
Canada	2,728562	Under Examination	2029
Europe***	09798248.2	Regional Phase - granted	
Denmark		Patent Registered	2029*
Finland		Patent Registered	2029*
France		Patent Registered	2029*
Germany		Patent Registered	2029*
		Patent Registered	2029*
Italy			
·		Patent Registered	2029*
Spain		Patent Registered Patent Registered	2029* 2029*
Spain Sweden			
Spain Sweden Switzerland The Netherlands		Patent Registered	2029*



Intellectual Property Report (continued...)

Europe ***	Divisional of 09798248.2	Filed	2029*
Japan	2011-516297	Under Examination	2029*
Japan	2014-208153 (Divisional of 2011-5516297)	Filed	2029*
USA	8,415,314	Patent Registered	2029*
USA	8,759,314	Patent Registered	2029*
ATL1102 Inhaled Ast	hma Patent Portfolio **		
International	PCT AU 2005/001634	National Phase applications	
Australia	AU 2005327506	Patent Registered	2025*
Canada	CA 2,584,614	Under examination	2025
Europe	EP1809302	Regional Phase - granted	
Denmark	DK/EP1809302T3	Patent Registered	2025*
Finland	EP(FI)1809302	Patent Registered	2025*
France	EP(FR)1809302	Patent Registered	2025*
Germany	DE 60 2005 035 821.8	Patent Registered	2025*
Italy	IT73129 BE/2012	Patent Registered	2025*
Spain	ES2392449	Patent Registered	2025*
Sweden	SE1809302T3	Patent Registered	2025*
United Kingdom	EP(UK)1809302	Patent Registered	2025*
Japan	JP 2007-535071	Abandoned	Relying on data exclusivity
New Zealand	NZ 554277	Patent Registered	2025
USA	US 8,765,700	Patent Registered	2028*
ATL1101 Patent Port	tfolio **		
International	PCT/AU2004/00160	National Phase applications	
Australia	2004210882	Patent Registered	2024 *
Canada	2515484	Patent Registered	2024
Europe	EP1597366	Regional Phase- granted	
Denmark	DK/EP1597366	Patent Registered	2024*
Finland	EP(FI)1597366	Patent Registered	2024*
France	EP(FR)1597366	Patent Registered	2024*
Germany	DE1597366	Patent Registered	2024*
Italy	IT1597366	Patent Registered	2024*
Spain	ES1597366	Patent Registered	2024*
Sweden	SE1597366	Patent Registered	2024*
United Kingdom	EP(UK)1597366	Patent Registered	2024*
Japan	4753863	Patent Registered	2024*
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Intellectual Property Report (continued...)

New Zealand	541637	Patent Registered	2024
USA	US7468356	Patent Registered	2025 *
USA	US8217017	Patent Registered	2025*
USA	9,084,770	Patent Registered	2029
USA	US14/731203 (continuation of US12/578,471)	Filed	2029

- * Potential for up to 5 year extensions to the patent term once the product is a registered drug.
- ** ATL1101, ATL1102, ATL1103 are also protected internationally by other Isis proprietary antisense technology patents and applications to which Antisense Therapeutics has world-wide license including US7015315 to 2023. Antisense technology patents are potentially extendible for up to 5 years to 2028 in the US.
- *** Designates all member states of European patent countries including all extension states.



Directors' Report

The Board of Directors of Antisense Therapeutics Limited present their report on the consolidated entity (referred to hereafter as 'the Company') consisting of Antisense Therapeutics Limited and the entities it controlled at the end of, or during, the year ended 30 June 2015. In order to comply with the provisions of the *Corporations Act 2001*, the Board of Directors report as follows:

Directors

The names of the Directors in office at any time during, or since the end of the year are as follows:

<u>'</u>	
Mr. Robert W Moses	Independent Non-Executive Chairman
Appointed to the Board	23 October 2001
Last elected by shareholders	1 November 2013
Qualifications	BA, MBA, FAICD, FAIM
Experience	Robert (Bob) Moses was formerly Corporate Vice President of CSL Limited. Mr. Moses draws on more than 40 years' experience in the pharmaceutical/biotechnology industry. During the period 1993-2001, Mr. Moses played a central role in CSL's development internationally. Prior to joining CSL, Mr. Moses was Managing Director of commercial law firm Freehills, Chairman and CEO of a NASDAQ listed medical service company, and Corporate Manager of New Business Development at ICI (now Orica). Mr. Moses is also the former Non-Executive Chairman of TGR Biosciences Pty Ltd. Mr. Moses also spent 17 years in various management roles at the multinational pharmaceutical company Eli Lilly.
Interest in shares and options	3,024,434 ordinary shares and 708,001 options over ordinary shares.
Committees	Chairman of the Remuneration Committee and member of the Audit Committee.
Directorships held in other listed entities	Nil
Mr. Mark Diamond	Managing Director

Mr. Mark Diamond	Managing Director
Appointed to the Board	31 October 2001
Qualifications	BSc, MBA, MAICD
Experience	Mark Diamond has over 25 years' experience in the pharmaceutical and biotechnology industry. Before joining Antisense Therapeutics Limited as MD and CEO in 2001, Mr. Diamond was employed in the US as Director, Project Planning/Business Development at Faulding Pharmaceuticals. Prior to this he held the positions of Senior Manager, Business Development and In-licensing within Faulding's European operation based in the UK and International Business Development Manager with Faulding in Australia.
Interest in shares and options	1,357,914 ordinary shares and 351,189 options over ordinary shares.
Committees	Nil
Directorships held in other listed entities	Nil



Dr. Chris Belyea	Independent Non-Executive Director
Appointed to the Board	13 November 2000
Last elected by shareholders	1 November 2013
Qualifications	BSc(Hons), PhD, FIPAA
Experience	Chris Belyea has a PhD in physics from the University of Melbourne and is a registered patent attorney. He became the founding CEO of Antisense Therapeutics Limited in November 2000 and remained in this role until January 2002 (shortly after Antisense Therapeutics Limited was listed on the Australian Stock Exchange). He worked for the Australian patent firm Griffith Hack & Co for 5 years before joining Circadian Technologies Limited as its Licensing and Projects Manager in 1996. In 1998 Dr. Belyea became founding CEO and member of the board of biotechnology company, Metabolic Pharmaceuticals Ltd. He served with Metabolic as an executive until mid-2008, and now runs his own patent attorney practice.
Interest in shares and options	285,579 ordinary shares and 61,222 options over ordinary shares.
Committees	Chairman of the Audit Committee and member of the Remuneration Committee.
Directorships held in other listed entities	Nil

Dr. Graham Mitchell	Independent Non-Executive Director
Appointed to the Board	24 October 2001
Last elected by shareholders	6 November 2014
Qualifications	AO, RDA, BVSc, FACVSc, PhD, FTSE, FAA
Experience	Graham Mitchell through Foursight Associates Pty Ltd, acts as joint Chief Scientist for the Victorian Government Department of Environment and Primary Industries. Dr. Mitchell is a Non-Executive Director of Avipep Pty Ltd and is a Principal of Foursight. Dr. Mitchell has held the position of Director of Research in the R&D Division of CSL Limited and for many years was a research scientist at The Walter & Eliza Hall Institute (WEHI). He is currently a Board Member of WEHI.
Interest in shares and options	240,180 ordinary shares and 60,582 options over ordinary shares.
Committees	Member of the Remuneration Committee.
Directorships held in other listed entities	Nil

Directors have been in office since the start of the financial year to the date of this report, unless stated otherwise.

Company Secretary

Mr. Phillip Hains held the position of Company Secretary since the start of the financial year to the date of this report.



Mr. Hains has served as the Company's Company Secretary and Chief Financial Officer since 9 November 2006. He is a Chartered Accountant operating a specialist public practice, 'The CFO Solution'.

The CFO Solution focuses on providing back office support, financial reporting and compliance systems for listed public companies. A specialist in the public company environment, Mr Hains has served the needs of a number of company boards and their related committees. He has over 20 years' experience in providing businesses with accounting, administration, compliance and general management services.

Principal Activity

The principal activity of Antisense Therapeutics Limited during the financial year was the research and development of novel antisense pharmaceuticals.

Dividends

The Directors did not pay any dividends during the financial year. The Directors do not recommend the payment of a dividend in respect of the 2015 financial year.

Significant Changes in State of Affairs

There have been no other significant changes in the nature of Antisense Therapeutics Limited's principal activities during the financial year.

Significant Events after Balance Date

There have not been any matters or circumstances, other than that referred to in the operations report, financial statements, or notes thereto, that have arisen since the end of the financial year, which significantly affected, or may significantly affect, the operations of Antisense Therapeutics Limited, the results of those operations or the state of affairs of Antisense Therapeutics Limited.

Likely Developments and Expected Results

The likely developments in the Company's operations, to the extent that such matters can be commented upon, are covered in the 'Operations Report'.

Operating and Financial Review

The profit of the Company after income tax for the financial year was \$706,918 (2014: loss \$3,013,272). This result has been achieved after fully expensing all research and development costs.

The Company had a cash reserve of \$6.8 million at 30 June 2015.

The 'Operations Report' provides further details regarding the progress made by the Company since the prior financial period, which have contributed to its results for the year.

Risk Management

The Board is responsible for overseeing the establishment and implementation of the risk management system, and to review and assess the effectiveness of the Company's implementation of that system on a regular basis.



The Board and senior management will continue to identify the general areas of risk and their impact on the activities of the Company. The potential risk areas for the Company include:

- efficacy, safety and regulatory risk of pre-clinical and clinical pharmaceutical development;
- financial position of the Company and the financial outlook;
- > economic outlook and share market activity;
- changing government policy (Australian and overseas);
- competitors' products/research and development programs;
- market demand and market prices for therapeutics;
- environmental regulations;
- > ethical issues relating to pharmaceutical research and development;
- the status of partnership and contractor relationships;
- other government regulations including those specifically relating to the biotechnology and health industries; and
- occupational health and safety and equal opportunity law.

Management will continue to perform a regular review of the following:

- the major risks that occur within the business;
- the degree of risk involved;
- the current approach to managing the risk; and
- where appropriate, determine:
 - o any inadequacies of the current approach; and
 - o possible new approaches that more efficiently and effectively address the risk.

Biotechnology Companies - Inherent Risks

Pharmaceutical research and development (R&D)

Pharmaceutical R&D involves scientific uncertainty and long lead times. Risks inherent in these activities include uncertainty of the outcome of the Company's research results; difficulties or delays in development of any of the Company's drug candidates; and general uncertainty related to the scientific development of a new medical therapy.

The Company's drug compounds require significant pre-clinical and human clinical development prior to commercialisation, which is uncertain, expensive and time consuming. There may be adverse side effects or inadequate therapeutic efficacy of the Company's drug candidates which would prevent further commercialisation. There may be difficulties or delays in testing any of the Company's drug candidates. There may also be adverse outcomes with the broader clinical application of the antisense technology platform which could have a negative impact on the Company's specific drug development and commercialisation plans.

No assurance can be given that the Company's product development efforts will be successful, that any potential product will be safe and efficacious, that required regulatory approvals will be obtained, that the Company's products will be capable of being produced in commercial quantities at an acceptable cost or at all, that the Company will have access to sufficient capital to successfully advance the products through development or to find suitable development or commercial partners for the development and or commercialisation of the products and that any products, if introduced, will achieve market acceptance.



Partnering and licensing

Due to the significant costs in drug discovery and development it is common for biotechnology companies to partner with larger biotechnology or pharmaceutical companies to help progress drug development. While the Company has previously entered into such licensing agreements with pharmaceutical partners, there is no guarantee that the Company will be able to maintain such partnerships or license its products in the future. There is also no guarantee that the Company will receive back all the data generated by or related intellectual property from its licensing partners. In the event that the Company does license or partner the drugs in its pipeline, there is no assurance as to the attractiveness of the commercial terms nor any guarantee that the agreements will generate a material commercial return for the Company.

Regulatory Approvals

Complex government health regulations, which are subject to change, add uncertainty to obtaining approval to undertake clinical development and obtain marketing approval for pharmaceutical products.

Delays may be experienced in obtaining such approvals, or the regulatory authorities may require repeat of different or expanded animal safety studies or human clinical trials, and these may add to the development cost and delay products from moving into the next phase of drug development and up to the point of entering the market place. This may adversely affect the competitive position of products and the financial value of the drug candidates to the Company.

There can be no assurance that regulatory clearance will be obtained for a product or that the data obtained from clinical trials will not be subject to varying interpretations. There can be no assurance that the regulatory authorities will agree with the Company's assessment of future clinical trial results.

Competition

The Company will always remain subject to the material risk arising from the intense competition that exists in the pharmaceutical industry. A material risk therefore exists that one or more competitive products may be in human clinical development now or may enter into human clinical development in the future. Competitive products focusing on or directed at the same diseases or protein targets as those that the Company is working on may be developed by pharmaceutical companies or other antisense drug companies including Isis or any of its other collaboration partners or licensees. Such products could prove more efficacious, safer, more cost effective or more acceptable to patients than the Company product. It is possible that a competitor may be in that market place sooner than the Company and establish itself as the preferred product.

Technology and Intellectual Property Rights

Securing rights to technology and patents is an integral part of securing potential product value in the outcomes of pharmaceutical R&D. The Company's success depends, in part, on its ability to obtain patents, maintain trade secret protection and operate without infringing the proprietary rights of third parties. There can be no assurance that any patents which the Company may own, access or control will afford the Company commercially significant protection of its technology or its products or have commercial application, or that access to these patents will mean that the Company will be free to commercialise its drug candidates. The granting of a patent does not guarantee that the rights of others are not infringed or that competitors will not develop technology or products to avoid the Company's patented technology or try to invalidate the Company's patents, or that it will be commercially viable for the Company to defend against such potential actions of competitors.



Environmental Regulation and Performance

The Company is involved in pharmaceutical research and development, much of which is contracted out to third parties, and it is the Director's understanding that these activities do not create any significant/material environmental impact. To the best of the Company's knowledge, the scientific research activities undertaken by, or on behalf of, the Company are in full compliance with all prescribed environmental regulations.

Meetings of Directors

During the financial year, 10 meetings of Directors (including committees of Directors) were held. Attendances by each Director during the year were as follows:

	Board M	leetings	Committee Meetings					
			Audit		Remune	eration		
	No. eligible to attend	No. attended	No. eligible to attend	No. attended	No. eligible to attend	No. attended		
Mr Robert W Moses	10	10	2	2	-	-		
Mr Mark Diamond	10	10	2	2	-	-		
Dr Chris Belyea	10	10	2	2	-	-		
Dr Graham Mitchell	10	8	2	2	-	-		

As at the date of this report the Company had an Audit Committee and Remuneration Committee, with membership of the committees as follows:

	Audit Committee	Remuneration Committee
Chairman	Dr Chris Belyea	Mr Robert W Moses
Members	Mr Robert W Moses	Dr Chris Belyea; and Dr Graham Mitchell

Indemnification and Insurance of Directors and other Officers

Under the Company's constitution:

- (a) To the extent permitted by law and subject to the restrictions in section 199A and 199B of the Corporations Act 2001, the Company indemnifies every person who is or has been an officer of the Company against any liability (other than for legal costs) incurred by that person as an officer of the Company where the Company requested the officer to accept appointment as Director.
- (b) To the extent permitted by law and subject to the restrictions in sections 199A and 199B of the Corporations Act 2001, the Company indemnifies every person who is or has been an officer of the Company against reasonable legal costs incurred in defending an action for a liability incurred by that person as an officer of the Company.

The Company has insured its Directors, the Company Secretaries and executive officers for the financial year ended 30 June 2015. Under the Company's Directors' and Officers' Liability Insurance Policy, the Company cannot release to any third party or otherwise publish details of the nature of the liabilities insured by the policy or the amount of the premium. Accordingly, the Company relies on section 300(9) of the Corporations Act 2001 to exempt it from the requirement to disclose the nature of the liability insured against and the premium amount of the relevant policy.

The Company also has in place a Deed of Indemnity, Access and Insurance with each of the Directors. This Deed:

- (i) indemnifies the Director to the extent permitted by law and the Constitution against certain liabilities and legal costs incurred by the Director as an officer of any Group Company;
- (ii) requires the Company to maintain, and pay the premium for, a D&O Policy in respect of the Director; and
- (iii) provides the Director with access to particular papers and documents requested by the Director for a Permitted Purpose,

both during the time that the Director holds office and for a seven year period after the Director ceases to be an officer of any Group Company, on the terms and conditions contained in the Deed.

Indemnification of Auditors

To the extent permitted by law, the Company has agreed to indemnify its auditors, Ernst & Young Australia, as part of the terms of its audit engagement agreement against claims by third parties arising from the audit (for an unspecified amount). No payment has been made to indemnify Ernst & Young during or since the financial year.

Share Options on Issue as at the Date of this Report

The unissued ordinary shares of Antisense Therapeutics Limited under option as at the date of this report were:

Class	Date of Expiry	Exercise Price	No. Under Option
ANPO	31 January 2017	\$0.27	46,950,984
ANPAU	30 July 2018	\$0.00	72,000

Proceedings on Behalf of the Company

No person has applied to the Court under section 237 of the *Corporations Act 2001* for leave to bring proceedings on behalf of the Company, or to intervene in any proceedings to which the Company is a party, for the purpose of taking responsibility on behalf of the Company for all or part of those proceedings.

No proceedings have been brought or intervened in on behalf of the Company with leave of the Court under section 237 of the *Corporations Act 2001*.

Non-Audit Services

The following non-audit services were provided by the entity's auditor, Ernst & Young. The Directors are satisfied that the provision of non-audit services is compatible with the general standard of independence for auditors imposed by the *Corporations Act 2001*. The nature and scope of each type of non-audit service provided means that auditor independence was not compromised.

Ernst & Young received or are due to receive the following amounts for the provision of non-audit services:

	2015 \$	2014 \$
Taxation Services	17,000	18,500

Auditor's Independence Declaration

The Auditors Independence Declaration as required under section 307C of the Corporations Act 2001 for the year ended 30 June 2015 has been received and can be found in the 'Auditor's Independence Declaration' section of this Annual Report.



Corporate Governance

In recognising the need for the highest standards of corporate behaviour and accountability, the Directors of Antisense Therapeutics support and adhere to good corporate governance practices. The Company's Corporate Governance Statement is contained in the 'Corporate Governance Statement' section of this Annual Report.

Remuneration Report (Audited)

This Remuneration Report outlines the Director and Executive remuneration arrangements of the Company as required by the Corporations Act 2001 and its Regulations.

This report details the nature and amount of remuneration of each Director of Antisense Therapeutics Limited and all other Key Management Personnel.

For the purposes of this report, Key Management Personnel (KMP) are defined as those persons having authority and responsibility for planning, directing and controlling the major activities of the Company, directly or indirectly, including any Director (whether Executive or otherwise) of the Company.

This report details the nature and amount of remuneration for each Director of Antisense Therapeutics Limited, and for the other Key Management Personnel.

The Directors of Antisense Therapeutics Limited during the year were:

Mr Robert W Moses Independent Non-Executive Chairman

Mr Mark Diamond Managing Director

Dr Chris Belyea Independent Non-Executive Director
Dr Graham Mitchell Independent Non-Executive Director

The other Key Management Personnel of Antisense Therapeutics Limited during the year were:

Dr George Tachas Director, Drug Discovery & Patents

Mr Phillip Hains Company Secretary and Chief Financial Officer

Section A: Principles used to determine the nature and amount of Remuneration Remuneration Policy

The Remuneration Policy ensures that Directors and Senior Management are appropriately remunerated having regard to their relevant experience, their performance, the performance of the Company, industry norms/standards and the general pay environment as appropriate. The Remuneration Policy has been established to enable the Company to attract, motivate and retain suitably qualified Directors and Senior Management who will create value for shareholders.

Remuneration Policy versus Company Performance

The Company's Remuneration Policy is not directly based on the Company's earnings. Prior to the year ended 30 June 2015, the Company's earnings had remained negative since inception due to the nature of the Company. Shareholder wealth reflects this speculative and volatile market sector. No dividends have ever been declared by the Company.

The Company continues to focus on the research and development of its intellectual property portfolio with the objective of achieving key development and commercial milestones in order to add further Shareholder value.

The Company's performance over the previous five financial years is as follows:

Net profit financial year 2015	\$706,918
Net loss financial year 2014	\$3,013,272
Net loss financial year 2013	\$2,454,842
Net loss financial year 2012	\$1,801,278
Net loss financial year 2011	\$1,813,550

The Company's share price over the previous five financial years is as follows:

30 June 2015	\$0.12
30 June 2014	\$0.14
30 June 2013	\$0.10
30 June 2012	\$0.18
30 June 2011	\$0.08

Remuneration Committee

The Remuneration Committee of the Board of Directors of Antisense Therapeutics Limited is responsible for overseeing the Remuneration Policy of the Company and for recommending or making such changes to the policy as it deems appropriate.

Non-Executive Director Remuneration

Objective

The Remuneration Policy ensures that Non-Executive Directors are appropriately remunerated having regard to their relevant experience, individual performance, the performance of the Company, industry norms/standards and the general pay environment as appropriate.

Structure

The Company's Constitution and the ASX Listing Rules specify that the aggregate remuneration of Non-Executive Directors shall be determined from time to time by a General Meeting. An amount (not exceeding the amount approved at the General Meeting) is determined by the Board and then divided between the Non-Executive Directors as agreed. The latest determination was at the General Meeting held on 13 November 2001 when shareholders approved the aggregate maximum sum to be paid or provided as remuneration to the Directors as a whole (other than the Managing Director and Executive Directors) for their services as \$300,000 per annum.

In the year ended 30 June 2015, the Non-Executive Directors were remunerated in aggregate \$130,293 per annum, excluding superannuation.

The manner in which the aggregate remuneration is apportioned amongst Non-Executive Directors is reviewed periodically.

The Board is responsible for reviewing its own performance. Board, and Board committee performance, is monitored on an informal basis throughout the year with a formal review conducted during the financial year.

No retirement benefits are payable other than statutory superannuation, if applicable.



Executive Director and Executive Officer Remuneration

Objective

The Remuneration Policy ensures that Executive Directors are appropriately remunerated having regard to their relevant experience, individual performance, the performance of the Company, industry norms/standards and the general pay environment as appropriate.

Structure

The Non-Executive Directors are responsible for evaluating the performance of the Managing Director, who in turn evaluates the performance of the other Senior Executives. The evaluation process is intended to assess the Company's business performance, whether long-term strategic objectives are being achieved and the achievement of individual performance objectives.

The performance of the Managing Director and Senior Executives are monitored on an informal basis throughout the year and a formal evaluation is performed annually.

Fixed Remuneration

Executives' fixed remuneration comprises salary and superannuation and is reviewed annually by the Managing Director, and in turn, the Remuneration Committee. This review takes into account the Executives' experience, performance in achieving agreed objectives and market factors as appropriate.

Variable Remuneration - Short Term Incentive Scheme

All Executives are entitled to participate in the Employee Short Term Incentive Scheme which provides for annual cash bonuses for outstanding performance in the achievement of key corporate and individual objectives. The Remuneration Committee approves the issue of cash bonuses following the recommendations of the Managing Director in his review of the performance of the Executives and the Company as a whole.

The Short Term Incentive Scheme operates as follows:

The Board determines whether Executives are eligible for bonuses on an annual basis. The cash bonuses, based on the recommendations of the Managing Director for outstanding performance, are not linked to any specific Key Result Areas (KRA's). The maximum achievable bonus for an Executive is 35% of the Executive's base salary. There were no bonuses paid under the Short Term Incentive Scheme during the year.

Variable Remuneration – Long Term Incentive Scheme

Executives may also be provided with longer-term incentives through the Company's Employee Option Plan, to allow the Executives to participate in and benefit from the growth of the Company as a result of their efforts and to assist in motivating and retaining those key employees over the long term. Continued service is the condition attached to the vesting of the options. The Board at its discretion determines the total number of options granted to each Executive. There were no options granted under the Long Term Incentive Scheme during the year.

Section B: Details of Remuneration

Details of Remuneration for the year ended 30 June 2015

The remuneration for each Director and each of the other Key Management Personnel of the Company during the year ended 30 June 2015 was as follows:

	Short-term employee benefits	Post-employment Benefits	Long-term Benefits	Total
30 Jun 2015	Cash salary and fees	Pension and Super Contribution \$	Long Service Leave \$	\$
Directors				
Mr Robert W Moses	56,293	5,348	-	61,641
Mr Mark Diamond	366,000	27,450	7,146	400,596
Dr Chris Belyea	37,500	3,563	-	41,063
Dr Graham Mitchell	36,500	3,468	-	39,968
	496,293	39,829	7,146	543,268
Other Key Management Personnel				
Dr George Tachas	220,185	20,918	4,300	245,403
Mr Phillip Hains ¹	99,000	-	-	99,000
	319,185	20,918	4,300	344,403
	815,478	60,747	11,446	887,671

Remunerated through The CFO Solution (see Section D below and the Company Secretary details above for further detail)

Details of Remuneration for the year ended 30 June 2014

The remuneration for each Director and each of the other Key Management Personnel of the Company during the year ended 30 June 2014 was as follows:



30 Jun 2014	Short-term employee benefits Cash salary and fees	Post-employment Benefits Pension and Super Contribution \$	Long-term Benefits Long Service Leave \$	Total \$
Directors				
Mr Robert W Moses	56,293	5,207	-	61,500
Mr Mark Diamond	366,000	27,450	7,157	400,607
Dr Chris Belyea	37,500	3,469	-	40,969
Dr Graham Mitchell	36,500	3,376	-	39,876
	496,293	39,502	7,157	542,952
Other Key Management Personnel				
Dr George Tachas	220,185	20,367	4,306	244,858
Mr Phillip Hains ¹	99,000	-	-	99,000
	319,185	20,367	4,306	343,858
	815,478	59,869	11,463	886,810

Remunerated through The CFO Solution (see Section D below and the Company Secretary details above for further detail)

Performance based Remuneration for the year ended 30 June 2015

	% of Total Remuneration for the Year	Estimated maximum value of	Estimated minimum value of	% of remuneration that is	% of remuneration that is non-
	that consisted of cash	bonus for the year	bonus for the year	performance based	performance based
30 June 2015	bonuses %	\$	\$	%	%
Directors					
Mr Robert W Moses	-	-	-	-	100%
Mr Mark Diamond	-	-	-	-	100%
Dr Chris Belyea	-	-	-	-	100%
Dr Graham Mitchell	-	-	-	-	100%
Other Key Management					
Personnel					
Dr George Tachas	-	-	-	-	100%
Mr Phillip Hains	-	-	-	-	100%



Performance based Remuneration for the year ended 30 June 2014

30 June 2014	% of Total Remuneration for the Year that consisted of cash bonuses %	Estimated maximum value of bonus for the year \$	Estimated minimum value of bonus for the year \$	% of remuneration that is performance based	% of remuneration that is non-performance based
Directors	76	· · · ·	· · ·	70	76
Mr Robert W Moses	_	_	_	_	100%
Mr Mark Diamond	_	35%	_	_	100%
Dr Chris Belyea		3370	_	_	100%
Dr Graham Mitchell	_	_	_	_	100%
Dr Graffalli Milleffeli	-	-	-	-	100%
Other Key Management					
Personnel					
Dr George Tachas	-	35%	-	-	100%
Mr Phillip Hains	-	-	-	-	100%

Section C: Share-based Compensation

(a) Shareholdings

The number of shares in the Company held during the financial year by each Director and other Key Management Personnel of the Company, including their personally related parties, are set out below.

No shares granted to Directors and Key Management Personal during the period as compensation.

30 June 2015	Balance at start of the year	Granted as Compensation	Options Exercised	Net Change Other	Balance at end of the year	Balance held nominally at the end of the reporting period
Directors						
Mr Robert W Moses	2,124,000	-	-	900,434	3,024,434	-
Mr Mark Diamond	1,053,567	-	-	304,347	1,357,914	-
Dr Chris Belyea	111,666	-	-	173,913	285,579	-
Dr Graham Mitchell	109,745	-	-	130,435	240,180	
	3,398,978	-	-	1,509,129	4,908,107	
Other Key Manageme	nt Personnel					
Dr George Tachas	485,324	-	-	173,912	659,236	-
Mr Phillip Hains	233,052	-	-	-	233,052	-
	718,376	-	-	173,912	892,288	-
	4,117,354	-	-	1,683,041	5,800,395	-



(b) Options and Rights

The number of options over ordinary shares in the Company held during the financial year by each Director of Antisense Therapeutics Limited and other Key Management Personnel of the Company, including their personally related parties, are set out below:

30 June 2015	Balance at start of the year	Granted as Compensation	Options Exercised	Net Change Other	Total vested at end of the year	Total Vested and exercisable at the end of the year	Total vested and unexercisable at the end of the year
<u>Directors</u>							
Mr Robert W Moses	708,001	-	-	-	708,001	708,001	-
Mr Mark Diamond	351,189	-	-	-	351,189	351,189	-
Dr Chris Belyea	61,222	-	-	-	61,222	61,222	-
Dr Graham Mitchell	60,582	-	-	-	60,582	60,582	-
	1,180,994	-	-	-	1,180,994	1,180,994	-
Other Key Management Personnel							
Dr George Tachas	159,276	-	-	-	159,276	159,276	-
Mr Phillip Hains	77,684	-	-	-	77,684	77,684	
	236,960	-	-	-	236,960	236,960	-
	1,417,954	-	-	-	1,417,954	1,417,954	-



Section D: Employment Contracts of Key Management Personnel

At the date of this report, the employment conditions of the Managing Director, Mr Mark Diamond and other Key Management Personnel were formalised in contracts of employment. Mr Mark Diamond is employed under a contract, which commenced on 31 October 2001. Subsequent to this contract a notice period for Mr Diamond of between two and four months was negotiated depending upon the party ending the agreement.

Antisense Therapeutics Limited has a contract with The CFO Solution, a specialist public practice, focusing on providing back office support, financial reporting and compliance systems for listed public companies. Through this contact the services of Mr Phillip Hains were provided. The contract commenced on 9 November 2006 and can be terminated with three months' notice of either party.

Section E: Additional Information

(a) Equity issued as part of remuneration for the year ended 30 June 2015

During the financial year ended 30 June 2015, no options were granted, exercised or lapsed by any of the Key Management Personnel.

(b) Loans to Directors and Other Key Management Personnel

There were no loans made to Directors or other Key Management Personnel of the Company, including their personally related parties.

(c) Other transactions with Other Key Management Personnel

Transactions between Key Management Personnel are on normal commercial terms and conditions no more favorable than those available to other parties unless otherwise stated. Transactions with related parties are as follows:

	2015	2014
	\$	\$
Purchases from Belyea IP		
Belyea IP is a patent attorney business operated by Dr Chris Belyea		
Service fees paid to Belyea IP during the year:	5,200	2,900
Patent renewals cost reimbursed to Belyea IP during the year:	36,422	28,793
Total paid by the Company to Belyea IP during the year:	41,622	31,693
At the end of the financial year, the Company owed Belyea IP:	-	-

This report is made in accordance with a resolution of Directors.



Mr Robert W Moses

Independent Non-Executive Chairman

Mr Mark Diamond **Managing Director**

Dated: This Day 21st of August 2015



Corporate Governance Statement

The Board of Directors of Antisense Therapeutics Limited ("the Company") is responsible for the corporate governance of the Company and guides and monitors the business and affairs of the Company on behalf of its shareholders.

The format of the Corporate Governance Statement is based on the Australian Stock Exchange Corporate Governance Council's ("the Council") "Corporate Governance Principles and Recommendations". In accordance with the Council's recommendations, the Corporate Governance Statement must contain certain specific information and must disclose the extent to which the Company has followed the guidelines during the period. Where a recommendation has not been followed, that fact must be disclosed, together will the reasons for the departure. The Company's Corporate Governance Statement is structured with reference to the Council's principles and recommendations, which are as follows:

Principle 1. Lay solid foundations for management and oversight

Principle 2. Structure the board to add value

Principle 3. Act ethically and responsibly

Principle 4. Safeguard integrity in corporate reporting

Principle 5. Make timely and balanced disclosure

Principle 6. Respect the rights of shareholders

Principle 7. Recognise and manage risk

Principle 8. Remunerate fairly and responsibly

Commensurate with the spirit of the ASX Corporate Governance Principles and Recommendations, the Company has followed each recommendation where the Board has considered the recommendation to be an appropriate benchmark for corporate governance practices, taking into account factors such as the size of the Company and the Board, resources available and activities of the Company. Where the Company's corporate governance practices depart from the Principles and Recommendations, the Board has offered full disclosure of the nature of, and reason for, the adoption of its own practice.

The Company's corporate governance practices were in place throughout the year ended 30 June 2015. For further information on the corporate governance policies adopted by the Company, please refer to its website: www.antisense.com.au.

Principle 1: Lay solid foundations for management and oversight.

Role of the Board

It is the role of the Board of Directors to represent and protect the interests of the Company's shareholders. The Board is responsible for the corporate governance of the Company and guides and monitors the business and affairs of the Company.

In furtherance of its responsibilities, the Board of Directors will:

- review, evaluate, provide input into and approve, on a regular basis, the Company's corporate governance strategy;
- monitor senior management's performance and implementation of strategy, and ensure appropriate resources are available;
- review, evaluate and approve the Company's budget and forecasts;
- review, evaluate, approve and monitor major resource allocations and capital investments, and any acquisitions and divestitures;
- review and monitor the financial and operating results of the Company;



- review and evaluate the overall corporate organisational structure, the assignment of senior management responsibilities and plans for senior management development and succession;
- > review, evaluate and approve compensation strategy as it relates to senior management of the Company;
- review and ratify systems of risk management and internal compliance and control, codes of conduct, and legal compliance;
- appoint and remove the Managing Director (Chief Executive Officer);
- ratify the appointment and, where appropriate, the removal of the Chief Financial Officer and the Company Secretary;
- > monitor its own performance and recommend and implement appropriate changes in composition and size.

Role of Management.

Through the Chief Executive Officer / Managing Director, management is responsible to the Board for the:

- 1) Development and implementation of agreed corporate strategy and performance objectives;
- 2) Undertaking the day to day activities of the Company;
- 3) Identifying all matters to be included in a risk profile of the Company and ensuring that effective risk management systems are implemented and adhered to;
- 4) Observing the code of conduct;
- 5) Ensuring that the Board is fully informed of all matters which may have a material impact on the ability of the Company to meet its obligations.

Board Appointments

The Company undertakes comprehensive reference checks prior to appointing a director, or putting that person forward as a candidate to ensure that person is competent, experienced, and would not be impaired in any way from undertaking the duties of director. The Company provides relevant information to shareholders for their consideration about the attributes of candidates together with whether the Board supports the appointment or re-election.

The terms of the appointment of a non-executive director, executive directors and senior executives are agreed upon and set out in writing at the time of appointment.

The Company Secretary

The Company Secretary is accountable directly to the Board, through the Chairman, on all matters to do with the proper functioning of the Board, including agendas, Board papers and minutes, advising the Board and its Committees (as applicable) on governance matters, monitoring that the Board and Committee policies and procedures are followed, communication with regulatory bodies and the ASX and statutory and other filings.

Diversity

The Company values the differences between its personnel and the valuable contribution that these differences can make to the Company. The Company is an equal opportunity employer and aims to recruit executives and employees from as diverse a pool of qualified candidates as reasonably possible based on their skills, qualifications and experience.

The Company is committed to increasing diversity amongst its employees, and not just in the area of gender diversity. Our workforce is employed based on the right person for the job regardless of their gender, age, nationality, race, religious beliefs, cultural background, sexuality or physical ability or appearance.



Executive and Board positions are filled by the best candidates available without discrimination. The Company is committed to increasing gender diversity within these positions when appropriate appointments become available. The Company is also committed to identifying suitable persons within the organisation, and where appropriate opportunities exist, advance diversity to support the promotion of talented employees into management positions.

The Company has not set any gender specific diversity objectives as it believes that multicultural diversity and other diversity factors are equally important within its organisation.

The following table demonstrates the Company's gender diversity as at 30 June 2015:

	Number of Males	Number of Females
Directors	4	-
Key Management Personnel	2	-
Other Company Employees	-	2

The Company employed 8 employees at the end of 30 June 2015 (2014: 8 employees).

Board Performance Review

The Board considers the ongoing development and improvement of its own performance, the performance of individual directors and Board Committees as critical to effective governance.

The Board has adopted an informal self-evaluation process to measure its own performance. The performance of the Board and individual directors is reviewed at least every year by the Board as a whole. This process includes a review in relation to the composition and skills mix of the Directors of the Company. Performance reviews involve analysis based on key performance indicators aligned with the financial and non-financial objectives of the Company. A performance review in accordance with the processes disclosed occurred during the 2015 financial year.

<u>Performance Review of KMP</u>

On at least an annual basis, the Board conducts a formal performance review of the Chief Executive Officer and any other key management personnel (KMP). The Board assesses the performance of KMP against qualitative and quantitative key performance indicators relevant to each KMP. A performance review of KMP occurred during the 2015 financial year in accordance with this process.

Independent Advice

The Board has procedures to allow Directors, in the furtherance of their duties, to seek independent professional advice at the Company's expense.

Principle 2: Structure the Board to add value.

Board composition

The length of service, skills, experience and expertise of each Director in office at the date of this report and throughout the 2015 financial year are included in the Directors' Report under the section headed 'Directors'. The Company's Board Charter stipulates that at least 50% of the Directors on the board should be independent Directors. Directors of Antisense Therapeutics Limited are considered to be independent when they are independent of management and free from any business or other relationship that could materially interfere with the exercise of their independent judgement.

In the context of Director independence, to be considered independent, a Non-Executive Director may not have a direct or indirect material relationship with the Company. The board considers that a material relationship is



one which impairs or inhibits, or has the potential to impair or inhibit, a Director's exercise of judgment on behalf of the Company and its shareholders.

From a quantitative perspective, an item is considered to be quantitatively immaterial if it is equal to or less than 5% of the relevant base amount. It is considered to be material (unless there is qualitative evidence to the contrary) if it is equal to or greater than 10% of the relevant base amount.

In accordance with the definition of independence above, and the materiality thresholds described, the majority of Directors are independent as set out below:

<u>Name</u>	<u>Position</u>
Mr Robert W Moses	Independent Non-Executive Chairman
Dr Graham Mitchell	Independent Non-Executive Director
Dr Chris Belyea	Independent Non-Executive Director

The term in office of each current Director is as follows:

<u>Name</u>	Term in Office	
Mr Robert W Moses	14 years	
Mr Mark Diamond	14 years	
Dr Chris Belyea	15 years	
Dr Graham Mitchell	14 years	

To ensure the Board is appropriately equipped to discharge its responsibilities, it has developed guidelines for the nomination and selection of Directors and for the operation of the Board. As the Antisense Therapeutics Limited's Board is not a large board, a formal nomination committee has not been established, as it is perceived that no real efficiencies would be gained from the existence of such a committee. The charter of the nomination committee has been incorporated into the Board Charter and by this action the Board of Directors considers all matters that would be relevant for a nomination committee. For additional details please refer to the Company's Board Charter on its website.

<u>Induction of New Directors and Ongoing Development</u>

Any new Directors will be issued with a formal Letter of Appointment that sets out the key terms and conditions of their appointment, including Director's duties, rights and responsibilities, the time commitment envisaged, and the Board's expectations regarding involvement with any Committee work.

A new director induction program is in place and Directors are encouraged to engage in professional development activities to develop and maintain the skills and knowledge needed to perform their role as Directors effectively.

Principle 3: Act ethically and responsibly

Code of Conduct

As part of its commitment to recognising the legitimate interests of stakeholders, the Company has established a Code of Conduct to guide compliance with legal and other obligations to legitimate stakeholders.

The Board acknowledges the legitimate interest of various stakeholders such as employees, clients, customers, government authorities, creditors and the community as a whole. As a good corporate citizen, it encourages compliance and commitment to appropriate corporate practices that are fair and ethical via its 'Code of Conduct'.

Trading in Company Securities.



The Company has a 'Code of Practice - Buying & Selling of Shares' that regulates the dealings by Directors and employees, in shares, options and other securities issued by the Company. The policy has been formulated to ensure that Directors and employees are aware of the legal restrictions on trading in Company securities while in possession of unpublished price sensitive information.

Principle 4: Safeguard integrity in corporate reporting.

Audit Committee

The Audit Committee operates under a charter approved by the Board. It is the Board's responsibility to ensure that an effective control framework exists within the entity. This includes ensuring that there are internal controls to deal with both the effectiveness and efficiency of significant business processes. This includes the safeguarding of assets, the maintenance of proper accounting records and the reliability of financial information as well as non-financial considerations. The Board has delegated the responsibility for the establishment and maintenance of a framework of internal control and ethical standards for the management of the Company to the Audit Committee.

The Audit Committee also provides the Board with additional assurance regarding the reliability of financial information for inclusion in the financial statements. All members of the Audit Committee are Non-Executive Directors. The Audit Committee is also responsible for the nomination of the external auditor and for reviewing the adequacy of the scope and quality of the annual statutory audit and half year statutory review. The Audit Committee Charter can be found on the Company's website.

The Audit Committee consists of two independent Non-Executive Directors. Given the current size of the Company, the Board believes that an Audit Committee consisting of two members is sufficient to enable the committee to discharge its mandate effectively. The members of the Audit Committee during the year were Dr Chris Belyea (Chairperson) and Mr Robert W Moses.

For details on the number of meetings for the Audit Committee held during the year and the attendances at those meetings, refer to the Directors' Report under the section headed 'Meetings of Directors'.

CEO and CFO Declarations

The CEO and CFO have provided the Board with a declaration that, in their opinion, the financial records of the entity have been properly maintained and that the financial statements comply with the appropriate accounting standards and give a true and fair view of the financial position and performance of the entity and that the opinion has been formed on the basis of a sound system of risk management and internal control which is operating effectively.

External Auditor

The Company's external auditor attends each annual general meeting and is available to answer any questions with regard to the conduct of the audit and their report.

Prior approval of the Board must be gained for non-audit work to be performed by the external auditor. There are qualitative limits on this non-audit work to ensure that the independence of the auditor is maintained.

There is also a requirement that the audit partner responsible for the audit not perform in that role for more than five years.

Principle 5: Making timely and balanced disclosure.

The Company has a Disclosure Policy which outlines the disclosure obligations of the Company as required under the ASX Listing Rules and Corporations Act. The policy is designed to ensure that procedures are in place so that



the market is properly informed of matters which may have a material impact on the price at which Company securities are traded.

The Board has designated the Company Secretary as the person responsible for overseeing and co-ordinating disclosure of information to the ASX as well as communicating with the ASX. In accordance with ASX Listing Rules the Company immediately notifies the ASX of information concerning the Company:

- (a) that a reasonable person would or may expect to have a material effect on the price or value of the Company's securities; and
- (b) that would, or would be likely to, influence persons who commonly invest in securities in deciding whether to acquire or dispose of the Company's securities.

Principle 6: Respect the rights of shareholders.

The Company is committed to providing current and relevant information to its shareholders.

The Company respects the rights of its shareholders, and to facilitate the effective exercise of the rights, the Company is committed to:

- (a) communicating effectively with shareholders through ongoing releases to the market via ASX information and general meetings of the Company;
- (b) giving shareholders ready access to balanced and understandable information about the Company and corporate proposals;
- (c) making it easy for shareholders to participate in general meetings of the Company; and

Any shareholder wishing to make inquiries of the Company is advised to contact the registered office. All public announcements made by the Company can be obtained from the ASX's website www.asx.com.au.

Shareholders may elect to, and are encouraged to, receive communications from the Company and its securities registry electronically.

The Company maintains information in relation to its corporate governance documents, Directors and senior executives, Board and committee charters, annual reports and ASX announcements on the Company's website.

Principle 7: Recognise and managing risk.

The Board is committed to the identification, assessment and management of risk throughout the Company's business activities.

The Board has established a policy for risk oversight and management within the Company. This is periodically reviewed and updated. Management reports risks identified to the Board through the monthly Operations Report, and via direct and timely communication to the Board where and when applicable. During the reporting period, management has reported to the Board as to the effectiveness of the Company's management of its material business risks. The Company does not have an internal audit function.

The Company faces risks inherent to its business, including economic risks, which may materially impact the Company's ability to create or preserve value for security holders over the short, medium or long term. The Company has in place policies and procedures, including a risk management framework (as described in the Company's Risk Management Policy), which is developed and updated to help manage these risks. The Board



does not consider that the Company currently has any material exposure to environmental or social sustainability risks.

The Company does not have separate risk committee. The Board as whole is responsible is responsible for overseeing the establishment and implementation of the risk management system. Due to the size of the Board and the Company, it is perceived that no real efficiencies would be gained from the existence of separate risk committee.

The Board review's the entity's risk management framework at least annually to satisfy itself that it continues to be sound. A review of the Company's risk management framework was conducted during the 2015 financial year.

Principle 8: Remunerate fairly and responsibly

It is the Company's objective to maintain a high quality Board and executive team by remunerating Directors at relevant market conditions. To assist in achieving this objective the Remuneration Committee remunerates Directors and executives having regard to their performance and the performance of the Company.

The expected outcomes of the remuneration policies and practices are to enable the Company to motivate, retain and attract Directors and executives who will create value for shareholders.

Details relating to the policy for performance evaluation and the amount of remuneration (monetary and non-monetary) paid to each Director and to each of the five highest-paid (non-director) executives during the year, are set out in the Directors' Report under the section headed 'Remuneration Report'.

The members of the Remuneration Committee at the date of this report were all independent Non-Executive Directors, being Mr Robert W Moses, Dr Chris Belyea and Dr Graham Mitchell. Details relating to performance evaluation are set out in the Directors' Report under the section headed 'Remuneration Report'. For details on the number of meetings of the Remuneration Committee held during the year and the attendees at those meetings, refer to the Directors' Report under the section headed 'Meetings of Directors'.

In accordance with the Company's share trading policy, participants in any equity based incentive scheme are prohibited from entering into any transaction that would have the effect of hedging or otherwise transferring the risk of any fluctuation in the value of any unvested entitlement in the Company's securities to any other person.

Further details in relation to the company's remuneration policies are contained in the Remuneration Report, within the Directors' report.



Auditor's Independence Declaration



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Auditor's Independence Declaration to the Directors of Antisense Therapeutics Limited

In relation to our audit of the financial report of Antisense Therapeutics Limited for the financial year ended 30 June 2015, to the best of my knowledge and belief, there have been no contraventions of the auditor independence requirements of the Corporations Act 2001 or any applicable code of professional conduct.

Ernst & Young

Frank & Young

Don Brumley Partner Melbourne

21 August 2015

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Annual Financial Statements For the year ended 30 June 2015

Statement of Profit or Loss and Other Comprehensive Income

For the year ended 30 June 2015

		Consolidated Entity		
		2015	2014	
	Note	\$	\$	
Revenue	3	3,916,337	82,936	
Other income	3	705,335	1,140,990	
Depreciation expenses	4	(8,172)	(9,753)	
Administration expenses	4	(1,884,169)	(1,830,981)	
Occupancy expenses	4	(115,397)	(115,238)	
Patent expenses	4	(205,353)	(153,477)	
Research and development expenses	4	(1,675,820)	(2,146,463)	
Foreign exchange gains/(losses)	4	(25,843)	18,714	
Profit/(loss) before income tax		706,918	(3,013,272)	
Income tax benefit	5	-		
Net profit/(loss) for the year		706,918	(3,013,272)	
Other comprehensive income/(loss) for the year				
Total comprehensive income/(loss) for the year		706,918	(3,013,272)	

		2015	2014
	Note	Cents	Cents
Earnings/(loss) per share for profit/(loss) attributable to the ordinary equity holders of the Company:			
Basic earnings/(loss) per share	8	0.45	(2.09)
Diluted earnings/(loss) per share	8	0.45	(2.09)

		Consolidat	ed Entity
		2015	2014
	Note	\$	\$
<u>ASSETS</u>			
<u>Current Assets</u>			
Cash and cash equivalents	9	6,829,605	1,334,513
Trade and other receivables	10	744,480	1,167,859
Prepayments		93,529	140,053
Total Current Assets		7,667,614	2,642,425
Non-Current Assets			
Plant and equipment	11	5,424	13,596
Total Non-Current Assets		5,424	13,596
TOTAL ASSETS		7,673,038	2,656,021
LIABILITIES			
LIABILITIES Command Machillaine			
Current Liabilities	12	204 004	240.001
Trade and other payables Borrowings	13 14	291,881	249,881 50,000
Provisions	15	289,559	269,249
Total Current Liabilities		581,440	569,130
			<u> </u>
TOTAL LIABILITIES		581,440	569,130
NET ASSETS		7,091,598	2,086,891
<u>EQUITY</u>			
Contributed equity	16	56,714,725	52,416,936
Reserves	17	960,855	960,855
Accumulated losses		(50,583,982)	(51,290,900)
TOTAL EQUITY		7,091,598	2,086,891

Statement of Changes in Equity

For the year ended 30 June 2015

Consolidated Entity	Contributed Equity	Option Reserve	Accumulated Losses	Total
Companies Emily	\$	\$	\$	\$
As at 30 June 2013	51,783,828	1,140,855	(48,277,628)	4,647,055
Profit/(loss) for the year	-	-	(3,013,272)	(3,013,272)
Total comprehensive income/(loss) for the year	-	-	(3,013,272)	(3,013,272)
Transactions with owners in their capacity as owners:				
Issue of shares	180,270	-	-	180,380
Options exercised net of costs	-	(180,000)	-	(180,000)
Options issued net of costs	454,378	-	-	454,378
Transaction costs on share issues	(1,540)	-	-	(1,540)
As at 30 June 2014	52,416,936	960,855	(51,290,900)	2,086,891
Profit/(loss) for the year	-	-	706,918	706,918
Total comprehensive income/(loss) for the year	-	-	706,918	706,918
Transactions with owners in their capacity as owners:				
Issue of shares	4,516,700	-	-	4,516,700
Transaction costs on share issues	(218,911)		-	(218,911)
As at 30 June 2015	56,714,725	960,855	(50,583,982)	7,091,598



		Consolidated Entity		
		2015	2014	
	Notes	\$	\$	
CASH FLOWS RELATED TO OPERATING ACTIVITIES				
Licensing fees received		3,863,988	-	
Payments to suppliers and employees		(3,775,898)	(4,167,717)	
Interest received		41,046	85,645	
R&D tax concession refund		1,139,739	974,187	
NET OPERATING CASH FLOWS	20a	1,268,875	(3,107,886)	
CASH FLOWS RELATED TO INVESTING ACTIVITIES			(10.615)	
Payment for purchases of plant and equipment NET INVESTING CASH FLOWS		<u> </u>	(10,615) (10,615)	
CASH FLOWS RELATED TO FINANCING ACTIVITIES				
Proceeds from issues of securities		4,445,128	563,696	
Capital raising costs		(218,911)	(110,588)	
NET FINANCING CASH FLOWS		4,226,217	453,108	
NET INCREASE/(DECREASE) IN CASH & CASH EQUIVALENTS		5,495,092	(2,665,393)	
Cash & cash equivalents at the beginning of the year		1,334,513	3,999,814	
Effects of exchange rate changes on cash & cash equivalents		-	92	
CASH & CASH EQUIVALENTS AT THE END OF THE YEAR	9,20	6,829,605	1,334,513	

Note 1. Statement of Significant Accounting Policies

Corporate Information

The financial report of Antisense Therapeutics Limited and its subsidiaries (the 'Company') for the year ended 30 June 2015 was authorised for issue in accordance with a resolution of the Directors on 21 August 2015. The financial report is for the Company consisting of Antisense Therapeutics Limited and its subsidiaries.

Antisense Therapeutics Limited is a listed public company limited by shares incorporated and domiciled in Australia whose shares are publicly traded on the Australian Securities Exchange. The Company also has a Level 1 ADR program traded on the US over-the-counter market.

The principal activity of the Company is the research and development of novel antisense pharmaceuticals.

Basis of Preparation

The financial report is a general-purpose financial report, which has been prepared in accordance with the requirements of the *Corporations Act 2001* and Australian Accounting Standards, required for a for-profit entity.

The financial report has been prepared on an accruals basis and is based on historical costs. The financial report is presented in Australian dollars, which is the Company's functional and presentation currency. All values are rounded to the nearest dollar unless otherwise stated.

Management is required to make judgements, estimates and assumptions about carrying values of assets and liabilities that are not readily apparent from other sources. The estimates and associated assumptions are based on historical experience and various other factors that are believed to be reasonable under the circumstance, the results of which form the basis of making the judgements. Actual results may differ from these estimates. The estimates and underlying assumptions are reviewed on an ongoing basis. Revisions to accounting estimates are recognised in the period in which the estimate is revised if the revision affects only that period, or in the period of the revision and future periods if the revision affects both current and future periods.

Judgements made by management in the application of Australian Accounting Standards that have significant effects on the financial statements and estimates with a significant risk of material adjustments in the next year are disclosed, where applicable, in the relevant notes to the financial statements.

Accounting policies are selected and applied in a manner which ensures that the resulting financial information satisfies the concepts of relevance and reliability, thereby ensuring that the substance of the underlying transactions or other events is reported.

Statement of Compliance

The financial report complies with Australian Accounting Standards as issued by the Australian Accounting Standards Board and International Financial Reporting Standards ("IFRS") as issued by the International Accounting Standards Board.

New, revised or amending Accounting Standards and Interpretations adopted

The consolidated entity has adopted all of the new, revised or amending Accounting Standards and Interpretations issued by the Australian Accounting Standards Board ('AASB') that are mandatory for the current reporting period.

Any new, revised or amending Accounting Standards or Interpretations that are not yet mandatory have not been early adopted. The following amending Standards have been adopted from 1 July 2014. Adoption of these Standards did not have any effect on the financial position or performance of the Company:

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Reference	Title	Summary
AASB 1031	Materiality	The revised AASB 1031 is an interim standard that cross-references to other Standards and the Framework (issued December 2013) that contain guidance on materiality. AASB 1031 will be withdrawn when references to AASB 1031 in all Standards and Interpretations have been removed.
		AASB 2014-1 Part C issued in June 2014 makes amendments to eight Australian Accounting Standards to delete their references to AASB 1031. The amendments are effective from 1 July 2014*.

Other than the amended accounting standards listed above, all other accounting standards adopted by the Company are consistent with the most recent Annual Report for the year ended 30 June 2014.

The following Australian Accounting Standards and Interpretations have recently been issued or amended but are not yet effective and therefore have not been adopted by the Company for the annual reporting period ended 30 June 2015:

Reference	Title	Summary	Application date of standard	Impact on financial report	Application date
AASB 9	Financial Instruments	AASB 9 addresses the classification, measurement and derecognition of financial assets and financial liabilities and introduces new rules for hedge accounting. In December 2014, the AASB made further changes to the classification and measurement rules and also introduced a new impairment model. These latest amendments now complete the new financial instruments standard.	1 January 2018	minimal	1 July 2018
AASB 15	Revenue from Contracts with Customers	The AASB has issued a new standard for the recognition of revenue. This will replace AASB 118 which covers contracts for goods and services and AASB 111 which covers construction contracts. The new standard is based on the principle that revenue is recognised when control of a good or service transfers to a customer – so the notion of control replaces the existing notion of risks and rewards. The standard permits a modified retrospective approach for the adoption. Under this approach entities will recognise transitional adjustments in retained earnings on the date of initial application (eg 1 July 2017), ie without restating the comparative period. They will only need to apply the new rules to contracts that are not completed as of the date of initial application The International Accounting Standards Board (IASB) in its July 2015 meeting decided to confirm its proposal to defer the effective date of IFRS 15 (the international equivalent of AASB 15) from 1 January 2017 to 1 January 2018. The amendment to give effect to the new effective date for IFRS 15 is expected to be issued in September 2015. At this time, it is expected that the AASB will make a corresponding amendment to AASB 15, which will mean that the application date of this standard for the Company will move from 1 July 2017 to 1 July 2018.	1 January 2017	Not yet assessed in detail	1 July 2017



Reference	Title	Summary	Application date of standard	Impact on financial report	Application date
AASB 2014-4	Clarification of Acceptable Methods of Depreciation and Amortisation (Amendments to AASB 116 and AASB 138)	AASB 116 Property Plant and Equipment and AASB 138 Intangible Assets both establish the principle for the basis of depreciation and amortisation as being the expected pattern of consumption of the future economic benefits of an asset. The IASB has clarified that the use of revenue-based methods to calculate the depreciation of an asset is not appropriate because revenue generated by an activity that includes the use of an asset generally reflects factors other than the consumption of the economic benefits embodied in the asset. The amendment also clarified that revenue is generally presumed to be an inappropriate basis for measuring the consumption of the economic benefits embodied in an intangible asset. This presumption, however, can be rebutted in certain limited circumstances.	1 January 2016	minimal	1 July 2016
AASB 2015-2	Amendments to Australian Accounting Standards – Disclosure Initiative: Amendments to AASB 101	The Standard makes amendments to AASB 101 Presentation of Financial Statements arising from the IASB's Disclosure Initiative project. The amendments are designed to further encourage companies to apply professional judgment in determining what information to disclose in the financial statements. For example, the amendments make clear that materiality applies to the whole of financial statements and that the inclusion of immaterial information can inhibit the usefulness of financial disclosures. The amendments also clarify that companies should use professional judgment in determining where and in what order information is presented in the financial disclosures.	1 January 2016	minimal	1 July 2016
AASB 2015-3	Amendments to Australian Accounting Standards arising from the Withdrawal of AASB 1031 Materiality	The Standard completes the AASB's project to remove Australian guidance on materiality from Australian Accounting Standards.	1 July 2015	minimal	1 July 2015

Accounting Policies

The following is a summary of the material accounting policies adopted by the Company in the preparation of the financial report. The accounting policies have been consistently applied, unless otherwise stated.

(a) Principles of Consolidation

The consolidated financial statements incorporate the assets and liabilities of all subsidiaries of Antisense Therapeutics Ltd as at 30 June 2015 and the results of all subsidiaries for the year then ended.

Subsidiaries are all those entities where the Company is exposed, or has rights, to variable returns from the Company's involvement with the entity and has the ability to affect those returns through the Company's power to direct the activities of the entity. The existence and effect of potential voting rights that are currently exercisable or convertible are considered when assessing whether the Company controls another entity.

Subsidiaries are fully consolidated from the date on which control is transferred to the Company. They are de-consolidated from the date that control ceases.

In preparing the consolidated financial statements, all intercompany balances and transactions, and unrealised profits/losses arising within the consolidated entity are eliminated in full. Investments in subsidiaries are accounted for at cost in the individual financial statements of Antisense Therapeutics Limited.



(b) Revenue recognition

Revenue is recognised to the extent that it is probable that the economic benefits will flow to the Company and the revenue can be reliably measured. The following specific recognition criteria must also be met before revenue is recognised.

Interest - control of the right to receive the interest payment.

Licensing revenue - right to receive the licensing revenue has been confirmed, and no significant obligations remain.

(c) Government grants

Government grants are recognised when there is reasonable assurance that the grant will be received and all grant conditions will be complied with.

When the grant relates to an expense item, it is recognised as income over the periods necessary to match the grant on a systematic basis to the costs that it is expected to compensate.

(d) Borrowing costs

Borrowing costs are expensed as incurred.

(e) Leases

The minimum lease payments of operating leases, where the lessor effectively retains substantially all of the risks and benefits of ownership of the leased item, are recognised as an expense on a straight-line basis.

(f) Cash and cash equivalents

Cash and short-term deposits in the Statement of Financial Position comprise cash at bank and in hand and short-term deposits with an original maturity of three months or less.

For the purposes of the Cash Flow Statement, cash and cash equivalents consist of cash and cash equivalents as defined above.

(g) Trade and other receivables

Trade and other receivables are recognised initially at fair value and subsequently measured at amortised cost using the effective interest method, less an allowance for impairment, once they become over due by more than 60 days. A separate account records the impairment.

An allowance for a doubtful debt is made when there is objective evidence that the Company will not be able to collect the debts. The criteria used to determine that there is objective evidence that an impairment loss has occurred include whether the Financial Asset is past due and whether there is any other information regarding increased credit risk associated with the Financial Asset. Bad debts which are known to be uncollectible are written off when identified.

(h) Foreign currency translation

The functional currency of the Company is based on the primary economic environment in which the Company operates. The functional currency of the Company is Australia dollars.

Transactions in foreign currencies are converted to local currency at the rate of exchange at the date of the transaction.

Amounts payable to and by the Company outstanding at reporting date and denominated in foreign currencies have been converted to local currency using rates prevailing at the end of the financial year.

All exchange differences are taken to profit or loss.

(i) Income tax

Deferred income tax is provided on all temporary differences at the balance date between the tax bases of assets and liabilities and their carrying amounts for financial reporting purposes.



Deferred income tax liabilities are recognised for all taxable temporary differences except where the deferred income tax liability arises from the initial recognition of an asset or liability in a transaction that is not a business combination and, at the time of the transaction, affects neither the accounting loss nor taxable profit or loss.

Deferred income tax assets are recognised for all deductible temporary differences, carry-forward of unused tax assets and unused tax losses, to the extent that it is probable that taxable profit will be available against which the deductible temporary differences, and the carry-forward of unused tax assets and unused tax losses can be utilised except where the deferred income tax asset relating to the deductible temporary differences arises from the initial recognition of an asset or liability in a transaction that is not a business combination and, at the time of transaction, affects neither the accounting loss nor taxable profit or loss.

The carrying amount of deferred income tax assets is reviewed at each balance date and reduced to the extent that it is no longer probable that sufficient taxable profit will be available to allow all or part of the deferred income tax asset to be utilised.

Deferred income tax assets and liabilities are measured at the tax rates that are expected to apply to the year when the asset is realised or the liability is settled, based on tax rates (and tax laws) that have been enacted or substantively enacted at balance date.

Deferred Tax assets are recognised for unused tax losses to the extent that it is probable that taxable profit will be available against which the losses can be utilised. Significant management judgement is required to determine the amount of deferred tax assets that can be recognised, based upon the likely timing and the level of future taxable profits together with future tax planning strategies.

Antisense Therapeutics Limited have not assessed unusued tax losses carried forward at 30 June 2015, given the history of losses from prior periods. These losses do not expire and may be used to offset taxable income in the current year and in future periods. Given the history of losses, there is limited support for the recognition of these losses as deferred tax assets. On this basis, Antisense Therapeutics Limited has determined it cannot recognise deferred tax assets on the tax losses carried forward. Further, on this basis, deferred tax assets have not been recognised related to temporary differences.

Income taxes relating to items recognised directly in equity are recognised in equity and not in profit or loss.

(j) Goods and services tax (GST)

Revenues, expenses and assets are recognised net of the amount of GST, except:

- where the GST incurred on a purchase of goods and services is not recoverable from the taxation authority, in which case the GST is recognised as part of the cost of acquisition of the asset or as part of the expense item as applicable; and
- receivables and payables are stated with the amount of GST included.

Cash flows arising from operating activities are included in the Cash Flow Statement on a gross basis (i.e. including GST) and the GST component of cash flows arising from investing and financing activities, which is recoverable from, or payable to, the taxation authority are classified as operating cash flows. Commitments and contingencies are disclosed net of the amount of GST recoverable from, or payable to, the taxation authority. The net amount of GST recoverable from or payable to, the taxation authority is included as part of the receivables or payables in the Statement of Financial Position.

(k) Plant and Equipment

Plant and equipment are measured at cost less any accumulated depreciation and any impairment losses. Such assets are depreciated over their useful economic lives as follows:

 Life	Method



Plant and equipment

3-5years

Straight line

(I) Intangible assets

Intangible assets are initially measured at cost. Following initial recognition, intangible assets are carried at cost less any accumulated amortisation and any accumulated impairment losses. The useful lives of intangible assets are assessed to be either finite or infinite. Intangible assets with finite lives are amortised over the useful life and assessed for impairment whenever there is an indication that the intangible asset may be impaired. The amortisation period and the amortisation method for an intangible asset with a finite useful life is reviewed at least at each financial year end. Changes in the expected useful life or the expected pattern of consumption of future economic benefits embodied in the asset are accounted for by changing the amortisation period or method, as appropriate, which is a change in an accounting estimate. The amortisation expense on intangible assets with finite lives is recognised in profit or loss in the expense category consistent with the function of the intangible asset.

(m) Research and Development Costs

Research costs are expensed as incurred.

An intangible asset arising from development expenditure on an internal project is recognised only when the Company can demonstrate the technical feasibility of completing the intangible asset so that it will be available for use or sale, its intention to complete and its ability to use or sell the asset, how the asset will generate future economic benefits, the availability of resources to complete the development and the ability to measure reliably the expenditure attributable to the intangible asset during its development.

Following initial recognition of the development expenditure, the cost model is applied requiring the asset to be carried at cost less any accumulated amortisation and accumulated impairment losses. Any expenditure so capitalised is amortised over the period of expected benefits from the related project.

The carrying value of an intangible asset arising from development expenditure is tested for impairment annually when the asset is not available for use, or more frequently when an indication of impairment arises during the reporting period.

(n) Impairment of non-financial assets

The carrying values of non-financial assets are tested for impairment whenever events or changes in circumstances indicate that the carrying amount may not be recoverable.

An impairment loss is recognised for the amount by which the asset's carrying amount exceeds its recoverable amount. Recoverable amount is the higher of an asset's fair value less costs of disposal and value in use. For the purposes of assessing impairment, assets are grouped at the lowest levels for which there are separately identifiable cash inflows that are largely independent of the cash inflows from other assets or groups of assets (cash-generating units). Non-financial assets that suffer an impairment are tested for possible reversal of the impairment whenever events or changes in circumstances indicate that the impairment may have reversed.

An impairment exists when the carrying value of an asset exceeds its estimated recoverable amount. The asset is then written down to its recoverable amount.

(o) Trade and other payables

Trade and other payables are carried at amortised cost and represent liabilities for goods and services provided to the Company prior to the end of the financial year that are unpaid and arise when the Company becomes obliged to make future payments in respect of the purchase of these goods and services. Licensing fees are recognised as an expense when it is confirmed that they are payable by the Company.

(p) Employee benefits



Wages, salaries and annual leave

Liabilities for wages and salaries, including non-monetary benefits and annual leave payments expected to be settled within 12 months of the reporting date are recognised in other provisions in respect of employees' service up to the reporting date. They are measured at the amounts expected to be paid when the liabilities are settled.

Long Service Leave

The liability for long service leave is recognised for employee benefits and measured as the present value of expected future payments to be made in respect of services provided by employees up to the reporting date. Consideration is given to expected future wage and salary levels, experience of employee departures, and periods of service. Expected future payments are discounted using market yields at the reporting date on national corporate bonds with terms to maturity and currencies that match, as closely as possible, to the estimated future cash outflows.

(q) Contributed equity

Ordinary shares are classified as equity. Any transaction costs arising on the issue of ordinary shares are recognised directly in equity as a reduction (net of tax) of the share proceeds received.

(r) Earnings per share

Basic earnings per share is calculated as net gain attributable to members, adjusted to exclude costs of servicing equity (other than dividends), divided by the weighted average number of ordinary shares, adjusted for any bonus element.

Diluted earnings per share is calculated as net gain attributable to members, adjusted for:

- costs of servicing equity (other than dividends);
- the after tax effect of dividends and interest associated with dilutive potential ordinary shares that have been recognised as expenses;
- other non-discretionary changes in revenues or expenses during the period that would result from the dilution of potential ordinary shares; divided by the weighted average number of ordinary shares and dilutive potential ordinary shares, adjusted for any bonus element.

(s) Parent Information

The financial information for the parent entity, Antisense Therapeutics Limited, disclosed in Note 2 has been prepared on the same basis as the consolidated statements with the exception of investments in subsidiaries which are carried at costs less any impairment.

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Note 2. Parent Information

The following information has been extracted from the books and records of the parent entity and has been prepared in accordance with the accounting standards.

	Parent	Entity
	2015	2014
Statement of Financial Position	\$	\$
<u>ASSETS</u>		
<u>Current Assets</u>		
Total Current Assets	7,667,614	2,642,425
Total Non-Current Assets	5,424	13,597
TOTAL ASSETS	7,673,038	2,656,022
LIABILITIES		
<u>Current Liabilities</u>		
Total Current Liabilities	581,440	569,131
TOTAL LIABILITIES	581,440	569,131
NET ASSETS	7,091,598	2,086,891
EQUITY		
Contributed equity	56,714,725	52,416,936
Reserves	960,855	960,855
Accumulated losses	(50,583,982)	(51,290,900)
TOTAL EQUITY	7,091,598	2,086,891

	Parent I	Parent Entity		
	2015	2014		
Statement of Comprehensive Income	\$	\$		
Net profit/(loss) for the year	706,918	(3,013,272)		
Total comprehensive income/(loss) for the year	706,918	(3,013,272)		



Note 3. Revenue and other income

		2015	2014
		\$	\$
Revenue			
Interest from external parties		52,349	82,936
Licensing revenue	3(a)	3,863,988	-
Total Revenue		3,916,337	82,936
Other income			
Government grants – R&D Tax incentive	3(b)	705,335	1,140,990
Total Other income		705,335	1,140,990
Total Revenue & Other Income		4,621,672	1,223,926

^{3 (}a) Licence fee received from Cortendo Caymen Limited.

Note 4. Expenses

	2015	2014
	\$	\$
Administration expenses		
Compliance expenses	220,171	252,295
Office expenses	61,875	90,558
Corporate employee expenses	673,807	672,655
Business development expenses	928,316	815,473
Total Administration expenses	1,884,169	1,830,981
Occupancy expenses		
Rent	98,777	98,777
Other expenses	16,620	16,461
Total Occupancy expenses	115,397	115,238
Research and development expenses		
R&D ATL 1101	-	-
R&D ATL 1102	267,051	616,232
R&D ATL 1103	1,251,433	1,374,370
R&D staff costs	157,336	155,861
Total Research and development expenses	1,675,820	2,146,463
Patent expenses	205,353	153,477
Depreciation expenses	8,172	9,753
Foreign exchange (gains)/losses	25,843	(18,714)
Total Expenses	3,914,754	4,237,198



^{3 (}b) Government grants related to research and development Tax incentives.

Note 5. Income Tax Benefit

		\$	\$
(a)	The components of tax benefit comprise:		
	Current tax	-	-
	Deferred tax	-	-
	Withholding tax refund on income earned in foreign tax jurisdiction	-	-
		-	
(b)	The prima facie tax on profit/(loss) from ordinary activities before tax at 30% (2014: 30%) is as follows:	212,075	(903,982)
	Add tax effect of:		
	Entertainment	587	483
	Share based payments	-	-
		587	483
	Less tax effect of:		
	Research and development tax concession	485,831	759,826
	Non-assessable grant income	(211,601)	(342,297)
	Section 40-880 deductions	(73,824)	(60,689)
		200,406	356,840
	Realisation of tax (benefit)/losses not previously recognised	(413,068)	546,659
	Withholding tax paid / (refund) on income earned in foreign tax jurisdiction	-	-
	Income tax (benefit) attributable to the Company	-	-
	The applicable weighted average effective tax rates are as follows:	0%	0%
(c)	Deferred Tax Assets and Liabilities		
	Foreign Exchange	-	-
	Accruals	883	12,716
	Provision for Annual Leave & Long Service Leave	6,093	3,948
	Other	10,566	11,402
	Gross Deferred Tax Assets	17,542	28,066
	Foreign Exchange	(772)	9,977
	Accruals	-	-
	Other	-	-
	Gross Deferred Tax Liabilities	(772)	9,977
	Net Deferred Tax Asset / (Liability) not recognised	18,314	18,089

Tax Losses

Antisense Therapeutics Limited has unconfirmed, unrecouped tax losses in Australia which have not been brought to account. The ability to be able to recognise a deferred tax asset in respect of these tax losses will be



dependent upon the probability that future taxable profit will be available against which the unused tax losses can be utilised and the conditions for deductibility imposed by Australian tax authorities will be complied with.

Note 6. Key Management Personnel Compensation

The aggregate compensation made to Directors and other Key Management Personnel of the Company is set out below:

	2015	2014
	\$	\$
Short-term employee benefits	815,478	815,478
Post-employment benefits	60,747	59,869
Long-term benefits	11,446	11,463
	887,671	886,810

For more information on Key Management Personnel Compensation, please refer to Remuneration Report contained under Directors' Report.

Note 7. Auditor's Remuneration

		2015 \$	2014 \$
Remun	eration of the auditor of the Company, Ernst & Young for:		
_	auditing or reviewing the financial report	49,244	47,741
	taxation services	17,000	18,500
		66,244	66,241

Note 8. Earnings/Losses per Share

	2015	2014
Basic earnings/(losses) per share (cents)	0.45	2.09
Diluted earnings/(losses) per share (cents)	0.45	2.09
 a) Net profit/(earnings/(losses)) used in the calculation of basic and diluted earnings/(losses) per share 	\$706,918	(\$3,013,272)
 Weighted average number of ordinary shares outstanding during the period used in the calculation of basic earnings/(losses) per share 	157,859,146	144,094,081
c) Adjustments for calculation of diluted earnings/(losses) per		
share: - Options over ordinary shares	72,000	Nil
 d) Weighted average number of ordinary shares outstanding during the period used in the calculation of diluted earnings/(losses) per share 	157,931,146	144,094,081



There have been no other conversions to, call of, or subscriptions for ordinary shares, or issues of potential ordinary shares since the reporting date and before the completion of this financial report.

Note 9. Cash and Cash Equivalents

	2015	2014
	\$	\$
Cash at bank and on hand	329,605	129,215
Term deposits	6,500,000	1,205,298
	6,829,605	1,334,513

The interest rate on cash at bank at 30 June 2015 was 0.10%p.a. (2014: 2.35% p.a.). And the interest rates on term deposits at 30 June 2015 were 2.15% p.a. (2014: 3.15% p.a.) for 30 days and 2.65% p.a. for 90 days. The term deposits have maturity periods of 30 days and 90 days.

Note 10. Trade and Other Receivables

	2015	2014
	\$	\$
Interest receivable	12,579	1,276
Australian Tax Office receivable	13,608	4,832
Research and development tax concession receivable	705,336	1,139,739
Other receivables	12,957	22,012
	744,480	1,167,859

Note 11. Plant and Equipment

	2015	2014
	\$	\$
At cost	172,209	172,209
Accumulated depreciation	(166,785)	(158,613)
Net book value	5,424	13,596
Balance at the beginning of the year	13,596	12,734
Additions	-	10,615
Depreciation expense	(8,172)	(9,753)
Balance at the end of the year	5,424	13,596



Note 12. Intangibles

	2015	2014
	\$	\$
At cost	6,387,500	6,387,500
Accumulated impairment losses / amortisation	(6,387,500)	(6,387,500)
	-	-

The intangible assets have finite useful lives.

- (a) The intangible assets relate to certain rights granted to Antisense Therapeutics Limited by Isis Pharmaceuticals Inc. ('Isis') upon listing of the Company. The main features of the agreement are as follows:
 - Isis has granted Antisense Therapeutics Limited certain rights to use Isis technology (i.e. Isis' patented technology) to commercialise antisense drugs to a number of protein targets (i.e. a research licence for each protein target). A certain number of these research licences to protein targets are also extendible to commercialisation licences.
 - The agreements with Isis provide access to and assistance in expanding Antisense Therapeutics Limited's drug pipeline and also provide access to and assistance in the Company's development projects including an exclusive license to a multiple sclerosis drug in Isis' preclinical pipeline; access to Isis manufacturing for provision of bulk quantities of antisense compounds for clinical trials; and access to Isis' preclinical development services for a sufficient period to allow smooth technology transfer.
- (b) The intangible assets were amortised on a straight-line basis over the term of the rights granted, five years. At 30 June 2007, the intangible assets had been fully amortised.

Note 13. Trade and Other Payables

	2015	2014
	\$	\$
Trade payables	161,804	116,860
Accrued expenses	125,500	128,444
Other payables	4,577	4,577
	291,881	249,881



Note 14. Borrowings

Unrestricted access was available at the reporting date to the following lines of credit:

	2015	2014
	\$	\$
Total facilities:		
Bank loan ¹	-	730,000
Used at the reporting date:		
Bank loan	-	50,000
Unused at the reporting date:		
Bank loan	-	680,000

¹ The bank loan relates to the secured funding facility the Company entered into with the Macquarie Bank Limited during the 2014 financial year. The facility was terminated on 15 October 2014.

Note 15. Provisions

	2015	2014
	\$	\$
Current employee provisions	289,559	269,249
Non-Current employee provisions	-	-
	289,559	269,249

Note 16. Contributed Equity

		2015	2014
	Note	\$	\$
Ordinary fully paid shares	16(a)	55,505,680	51,207,891
Options over ordinary shares	16(b)	1,209,045	1,209,045
		56,714,725	52,416,936



Note 16. Contributed Equity (Continued)

16(a) Ordinary Shares	2015		2014	
10(a) Ordinary Shares	No.	\$	No.	\$
Balance at the beginning of the year	144,096,128	51,207,891	1,437,954,566	51,029,161
Shares issued during the year	32,416,355	4,516,700	3,001,000	180,270
Consolidation 10:1 Nov 2013	-	-	(1,296,859,438)	-
Transaction costs relating to share issues	-	(218,911)	-	(1,540)
Balance at the end of the year	176,512,483	55,505,680	144,096,128	51,207,891

2015	Details	Number	Issue Price	
			\$	\$
1 October 2014	Placement Share Purchase	7,913,043	0.1150	910,000
12 November 2014 15 May 2015	Plan Issue of shares to Cortendo Cayman	9,478,237	0.1150	1,090,000
•	Limited	15,025,075	0.1675	2,516,700
	Transaction costs			(218,911)
		32,416,355		4,297,789

2014	Details	Number	Issue Price	
			\$	\$
2 July 2013	Exercise of ANPAU Unlisted Options	3,000,000	-	180,000
13 November 2013	Consolidation of shares 10:1 basis Exercise of ANPO	(1,296,859,438)	-	-
8 January 2014	Options	1,000	0.270	270
	Transaction costs			(1,540)
		(1,293,858,438)		178,730

Ordinary shares participate in dividends and the proceeds on winding up of the Company in proportion to the number of shares held. At shareholder meetings each ordinary share is entitled to one vote when a poll is called, otherwise each shareholder has one vote on a show of hands. The ordinary shares have no par value.



Note 16. Contributed Equity (Continued)

16 (b) Options	2015		2014	
10 (b) Options	No.	\$	No.	\$
Balance at the beginning of the year	46,950,984	1,209,045	-	754,667
Options issued during the year	-	-	46,951,984	563,426
Options exercised during the year	-	-	(1,000)	(110)
Options expired during the year	-	-	-	-
Transaction costs relating to option issues	-	-	-	(108,938)
Balance at the end of the year	46,950,984	1,209,045	46,950,984	1,209,045

2015	Details	Number	Issue Price \$
There has been	no activity during the financial year		

2014	Details	Number	Issue Price \$
20 November 2013	Issue of Loyalty Options (ANPO)	45,728,528	548,982
20 November 2013	Issue of Private Placement Options Issue of Options for Management Fees to	223,456	2,444
20 November 2013	Patersons	1,000,000	12,000
8 January 2014	Exercise of Loyalty Options (ANPO)	(1,000)	(110)
Capital raising costs ass	ociated with the above issues	-	(108,938)
		46,950,984	454,378

Note 17. Reserves

(a) Nature and Purpose of the Reserve

The option reserve recognises the proceeds from the issue of options over ordinary shares and the expense recognised in respect of share based payments, see note 21 for further detail.



Note 17. Reserves (Continued)

	2015		201	.4
	No.	\$	No.	\$
Unlisted options over fully paid ordinary				_
shares	72,000	960,855	3,720,000	1,140,855
Options exercised	-	-	(3,000,000)	(180,000)
Consolidation on 10:1 basis Nov 2013	-	-	(648,000)	_
	72,000	960,855	72,000	960,855

2015	Details	Number	Issue Price \$
No changes during the period.			
		-	_

2014	Details	Number	Issue Price \$
2 July 2013	Exercise of ANPAU Unlisted Options	(3,000,000)	(180,000)
13 November 2013	Consolidation on 10:1 basis	(648,000)	
		(3,648,000)	(180,000)

Options Outstanding at 30 June 2015

	No. of Options		
Date of Issue	27 Oct 2008	20 Nov 2013	
On issue at beginning of year	72,000	46,950,984	
Issued during the year	-	-	
Exercised during the year	-	-	
Expired during the year	-	-	
Forfeited during the year	-	-	
Consolidation 10:1 Nov 2013	-	-	
Outstanding at balance date	72,000	46,950,984	
Expired subsequent to balance date	-	-	
Exercised subsequent to balance date	-	-	
Outstanding at date of Directors' Report	72,000	46,950,984	
Original number of recipients	4	849	
Number of current holders	4	818	
Exercise price	-	\$0.27	
Exercise period from	27 Oct 2008	20 Nov 2013	
To (expiration day)	30 Jul 2018	31 Jan 2017	
The following proportion of options vest from the dates shown:			
100%	27 Oct 2008	20 Nov 2013	



Note 18. Commitments and Contingencies

	2015	2014
	\$	\$
Lease expenditure commitments:		
- not later than 12 months	24,693	24,693
- between 12 months and 5 years	-	-
	24,693	24,693

The lease expenditure commitments relate to the leasing of office premises. The lease is for a term of one year, expiring October 2015.

Note 19. Operating Segments

The Company has identified its operating segments based on the internal reports that are reviewed and used by the management team in assessing performance and determining the allocation of resources.

The operating segments are identified by management based on the manner in which the expenses are incurred, and for the purpose of making decisions about resource allocation and performance assessment. Discrete financial information about each of these operating segments is reported by the executive management team to the board on a regular basis.

Segments:

- ATL 1102 Multiple Sclerosis
- > ATL 1103 Growth and Sight Disorders

30 June 2015	Note	ATL1102 Multiple Sclerosis	ATL1103 Growth and Sight Disorders	Total
<u>Revenue</u>				
Segment Revenue		-	3,863,988	3,863,988
Unallocated Revenue	19a	-	-	52,349
Total Revenue		-	3,863,988	3,916,337
<u>Result</u>				
Segment Result		(99,520)	(718,548)	(818,068)
Unallocated Result	19b	-	-	(2,391,351)
Income Tax Benefit		-	-	-
Net Result		(99,520)	3,145,440	706,918

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Note 19. Operating Segments (Continued)

30 June 2014	Note	ATL1102 Multiple Sclerosis	ATL1103 Growth and Sight Disorders	Total
<u>Revenue</u>				
Segment Revenue		-	-	-
Unallocated Revenue	19a	-	-	82,936
Total Revenue		-	-	82,936
<u>Result</u>				
Segment Result		(262,034)	(590,010)	(852,044)
Unallocated Result	19b	-	-	(2,244,164)
Income Tax Benefit		-	-	1
Net Result		(262,034)	(590,010)	(3,013,272)

		2015	2014
		\$	\$
19(a)	Unallocated Revenue		
	- Interest from external parties	52,349	82,936
		52,349	82,936
19(b)	Unallocated Result		
	- R&D Tax Concession Refund	4,919	2,432
	- Compliance expenses	(220,171)	(252,295)
	- Business development expenses	(928,316)	(815,473)
	- Employee expenses	(673,807)	(672,655)
	- Patent expenses	(205,353)	(153,477)
	- Other expenses	(368,623)	(352,696)
		(2,391,351)	(2,244,164)



Note 20. Cash Flow Information

(a) Reconciliation of cash flow from operations with loss after income tax

	2015 \$	2014 \$
Net profit for the year	706,918	(3,013,272)
Add back depreciation expense	8,172	9,753
Add back share based payments	71,572	-
(Increases) in trade and other receivables	423,379	(154,601)
(Increases)/Decreases in prepayments	46,524	35,297
Increases/(Decreases) in trade and other payables	42,000	(48,130)
Increases in other current liabilities	(50,000)	50,000
Increases in provisions	20,310	13,159
Add back foreign exchange	<u> </u>	(92)
Net cash flows used in operating activities	1,268,875	(3,107,886)

Note 21. Events after the Balance Date

There have not been any matters or circumstances, other than that referred to in the financial statements or notes thereto, that have arisen since the end of the financial year, which significantly affected, or may significantly affect, the operations of Antisense Therapeutics Limited, the results of those operations or the state of affairs of Antisense Therapeutics Limited in future financial years.

Note 22. Related Party Transactions

Transactions between related parties are on normal commercial terms and conditions no more favorable than those available to other parties unless otherwise stated. Transactions with related parties are as follows:

	\$	\$
Purchases from Belyea IP		
Belyea IP is a patent attorney business operated by Dr Chris Belyea		
Service fees paid to Belyea IP during the year:	5,200	2,900
Patent renewals cost reimbursed to Belyea IP during the year:	36,422	28,793
Total paid by the Company to Belyea IP during the year: At the end of the financial year, the Company owed Belyea IP:	41,622	31,693



Note 23. Financial Risk Management Objectives and Policies

(a) Financial Instruments

The Company's financial instruments consist of cash and cash equivalents, trade and other receivables, trade and other payables and borrowings:

	2015	2014
	\$	\$
Cash and cash equivalents	6,829,605	1,334,513
Trade and other receivables	744,480	1,167,859
Trade and other payables	291,881	249,881
Borrowings	-	50,000

The Company does not have any derivative instruments at 30 June 2015 (2014: Nil).

(b) Risk Management Policy

The Board is responsible for overseeing the establishment and implementation of the risk management system, and reviews and assesses the effectiveness of the Company's implementation of that system on a regular basis.

The Board and Senior Management identify the general areas of risk and their impact on the activities of the Company, with Management performing a regular review of:

- > the major risks that occur within the business;
- the degree of risk involved;
- the current approach to managing the risk; and
- > if appropriate, determine:
 - o any inadequacies of the current approach; and
 - o possible new approaches that more efficiently and effectively address the risk.

Management report risks identified to the Board through the monthly Operations Report.

The Company seeks to ensure that its exposure to undue risk which is likely to impact its financial performance, continued growth and survival is minimised in a cost effective manner.

(c) Significant Accounting Policy

Details of significant accounting policies and methods adopted, including the criteria for recognition, the basis for measurement and the basis on which income and expenses are recognised, in respect of each class of financial asset, financial liability and equity instrument are disclosed in Note 1 to the financial statements.

The carrying amounts of cash and cash equivalents, trade and other receivables and trade and other payables represents their fair values determined in accordance with the accounting policies disclosed in note 1.

Interest revenue on cash and cash equivalents and foreign exchange movements on trade and other receivables and trade and other payables are disclosed in notes 3 and 4.



Note 23. Financial Risk Management Objectives and Policies (continued)

(d) Capital Risk Management

The Company's objectives when managing capital are to safeguard the Company's ability to continue as a going concern and to maintain an optimal capital structure so as to maximise shareholder value. In order to maintain or achieve an optimal capital structure, the Company may issue new shares or reduce its capital, subject to the provisions of the Company's constitution. The capital structure of the Company consists of equity attributed to equity holders of the Company, comprising contributed equity, reserves and accumulated losses disclosed in notes 16 and 17. By monitoring undiscounted cash flow forecasts and actual cash flows provided to the Board by the Company's Management the Board monitors the need to raise additional equity from the equity markets.

(e) Financial Risk Management

The main risks the Company is exposed to through its operations are interest rate risk, foreign exchange risk, credit risk and liquidity risk.

Interest Rate Risk

The Company is exposed to interest rate risks via the cash and cash equivalents that it holds. Interest rate risk is the risk that a financial instruments value will fluctuate as a result of changes in market interest rates. The objective of managing interest rate risk is to minimise the Company's exposure to fluctuations in interest rate that might impact its interest revenue and cash flow.

To manage interest rate risk, the Company locks a portion of the Company's cash and cash equivalents into term deposits. The maturity of term deposits is determined based on the Company's cash flow forecast.

Interest rate risk is considered when placing funds on term deposits. The Company considers the reduced interest rate received by retaining cash and cash equivalents in the Company's operating account compared to placing funds into a term deposit. This consideration also takes into account the costs associated with breaking a term deposit should early access to cash and cash equivalents be required.

The Company's exposure to interest rate risk and the weighted average interest rates on the Company's financial assets and financial liabilities is as follows:



Note 23. Financial Risk Management Objectives and Policies (Continued)

30 June 2015	Weighted Average Effective Interest Rate	Floating Interest Rate	Fixed Interest Rate within Year	Fixed Interest Rate 1 to 5 years	Fixed Interest Rate over 5 Years	Non- Interest Bearing	Total
	%	\$	\$	\$	\$	\$	\$
Financial Assets Cash and cash equivalents Trade and other	2.53	329,205	6,500,000	-	-	400	6,829,605
receivables	_	_	_	_	_	744,480	744,480
		329,205	6,500,000	-	-	744,880	7,574,085
Financial Liabilities Trade and other payables	-	-	-	-	-	291,881 291,881	291,881 291,881
30 June 2014	Weighted Average Effective Interest Rate	Floating Interest Rate	Fixed Interest Rate within Year	Fixed Interest Rate 1 to 5 years	Fixed Interest Rate over 5 Years	Non- Interest Bearing	Total
				o yours			
	%	\$	\$	\$	\$	\$	\$
Financial Assets Cash and cash							
Cash and cash equivalents Trade and other	3.51	\$ 128,815	\$ 1,205,298			400	1,334,513
Cash and cash equivalents		128,815 -	1,205,298		\$ - -	400 1,167,859	1,334,513 1,167,859
Cash and cash equivalents Trade and other receivables Financial Liabilities Trade and other						400	1,334,513
Cash and cash equivalents Trade and other receivables Financial Liabilities		128,815 -	1,205,298		\$ - -	400 1,167,859 1,168,259	1,334,513 1,167,859 2,502,372

There has been no change to the Company's exposure to interest rate risk or the manner in which it manages and measures its risk in the year ended 30 June 2015.

The Company has conducted a sensitivity analysis of the Company's exposure to interest rate risk. The percentage change is based on the expected volatility of interest rates using market data and analysts forecasts. The analysis shows that if the Company's interest rate was to fluctuate as disclosed below and all other variables had remained constant, then the interest rate sensitivity impact on the Company's profit after tax and equity would be as follows:



Note 23. Financial Risk Management Objectives and Policies (Continued)

	(Higher) / Lower 2015	(Higher) / Lower 2014
2015: +1% (2014: +1%)	68,296	13,345
2015: - 1% (2014: -1%)	(68,296)	(13,345)

Foreign Currency Risk

The Company is exposed to foreign currency risk via the trade and other receivables and trade and other payables that it holds. Foreign currency risk is the risk that the value of a financial instrument will fluctuate due to changes in foreign exchange rates. The Company aims to take a conservative position in relation to foreign currency risk hedging when budgeting for overseas expenditure however; the Company does not have a policy to hedge overseas payments or receivables as they are highly variable in amount and timing, due to the reliance on activities carried out by overseas entities and their billing cycle.

The following financial assets and liabilities are subject to foreign currency risk:

	2015	2014
	\$	\$
Cash and cash equivalents (AUD/GBP)	-	1,029
Trade and other payables (AUD/USD)	31,109	40,362
Trade and other payables (AUD/GBP)	13,899	44,115
Trade and other payables (AUD/EUR)	10,108	2,329

Foreign currency risk is measured by regular review of our cash forecasts, monitoring the dollar amount and currencies that payment are anticipated to be paid in. The Company also considers the market fluctuations in relevant currencies to determine the level of exposure. If the level of exposure is considered by Management to be too high, then Management has authority to take steps to reduce the risk.

Steps to reduce risk may include the acquisition of foreign currency ahead of the anticipated due date of an invoice or may include negotiations with suppliers to make payment in our functional currency. Management mitigated foreign currency risk by purchasing Great British Pounds currency during the current financial year. Should Management determine that the Company should consider taking out a hedge to reduce the foreign currency risk, they would need to seek Board approval.

The Company conducts some activities outside of Australia which exposes it to transactional currency movements, where the Company is required to pay in a currency other than its functional currency.

There has been no change in the manner the Company manages and measures its risk in the year ended June 2015.

The Company is exposed to fluctuations in United States dollars, Euros, and Great British Pounds. Analysis is conducted on a currency by currency basis using sensitivity variables.

The Company has conducted a sensitivity analysis of the Company's exposure to foreign currency risk. The sensitivity analysis variable is based on the expected overall volatility of the significant currencies, which is based on management's assessment of reasonable possible fluctuations taking into consideration movements over the last 6 months each year and the spot rates at each reporting date. The analysis shows that if the Company's exposure to foreign currency risk was to fluctuate as disclosed below and all other variables had remained constant, then the foreign currency sensitivity impact on the Company's loss after tax and equity would be as follows:



Note 23. Financial Risk Management Objectives and Policies (Continued)

	(Higher) / Lower 2015	(Higher) / Lower 2014
Cash and cash equivalents		
AUD/GBP: 2015: +3% (2014: +3%)	-	31
AUD/GBP: 2015: -3% (2014: -3%)	-	(31)
Trade and other payables		
AUD/USD: 2015: +3% (2014: +3%)	(933)	(1,211)
AUD/USD: 2015: -3% (2014: -3%)	933	1,211
AUD/GBP: 2015: +3% (2014: +3%)	417	1,323
AUD/GBP: 2015: -3% (2014: -3%)	(417)	1,323
AUD/EUR: 2015: +3% (2014: +3%)	303	70
AUD/EUR: 2015: -3% (2014: -3%)	(303)	(70)

Credit Risk

The Company is exposed to credit risk via its cash and cash equivalents and trade and other receivables. Credit risk is the risk that a counter-party will default on its contractual obligations resulting in a financial loss to the Company. To reduce risk exposure for the Company's cash and cash equivalents, it places them with high credit quality financial institutions.

Historically the Company has had minimal trade and other receivables, with the majority of its funding being provided via shareholder investment. Traditionally the Company's trade and other receivables relate to GST refunds and Research and Development Tax Concession amounts due to the Company from the Australian Tax Office. At 30 June 2015 GST accounted for \$13,608 (2014: \$4,832) of the trade and other receivables, respectively. At 30 June 2015, accrued interest from the Commonwealth Bank amounted to \$12,579 (2014: \$911).

The trade and other receivables at 90+ days also include the rent bond on the office premises of \$8,231. This is not considered impaired. The Board believes that the Company does not have significant credit risk at this time in respect of its trade and other receivables.

The Company has analysed its trade and other receivables below. All trade and other receivables disclosed below have not been impaired.

	0-30 days \$	31-60 days \$	61-90 days \$	90+ days \$
2015 Trade and other receivables	736,249	-	-	8,231
2014 Trade and other receivables	1,159,628	-	-	8,231



Note 23. Financial Risk Management Objectives and Policies (Continued)

Liquidity Risk

The Company is exposed to liquidity risk via its trade and other payables. Liquidity risk is the risk that the Company will encounter difficulty in raising funds to meet the commitments associated with its financial instruments. Responsibility for liquidity risk rests with the Board who manage liquidity risk by monitoring undiscounted cash flow forecasts and actual cash flows provided to them by the Company's Management at Board meetings to ensure that the Company continues to be able to meet its debts as and when they fall due. Contracts are not entered into unless the Board believes that there is sufficient cash flow to fund the associated commitments. The Board considers when reviewing its undiscounted cash flow forecasts whether the Company needs to raise additional funding from the equity markets.

The Company has analysed its trade and other payables below:

	0-30 days \$	31-60 days \$	61-90 days \$	90+ days \$
2015 Trade and other payables	291,881	-	-	-
2014 Trade and other payables	249,881	-	-	-

Note 24. Subsidiaries

The consolidated financial statements incorporate the assets, liabilities and results of the following subsidiaries in accordance with the accounting policy described in note 1:

Name of entity	Country of incorporation	Percentage owned (%)	
		2015	2014
Downsk Fuelika			
Parent Entity			
Antisense Therapeutics Limited			
Subsidiaries of Antisense Therapeutics Limited			
Antisense Therapeutics (HK) Pty Ltd ¹	Australia	100	100

¹ On 10 July 2012 the parent entity incorporated Antisense Therapeutics (HK) Pty Ltd, a wholly owned subsidiary. The purpose of this new incorporated entity is to facilitate the provision of the relevant licenses to ATL1102 intellectual property in a proposed Joint Venture with a Chinese Company.

Note 25. Company Details

The registered office of the Company is:

6-8 Wallace Avenue, Toorak, Victoria, 3142

The principal place of business of the Company is:

6-8 Wallace Avenue, Toorak, Victoria, 3142



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Directors' Declaration

The Directors of the Company declare that:

In the opinion of the Directors:

- 1. the financial statements and notes, as set out on pages 35 to 64 are in accordance with the Corporations Act 2001 and:
 - a. comply with Accounting Standards and the Corporations Regulations 2001; and
 - b. give a true and fair view of the financial position as at 30 June 2015 and of the performance for the year ended on that date of the Company;
 - c. the financial statements and notes also comply with International Financial Reporting Standards as disclosed in Note 1.
- 2. in the Directors' opinion there are reasonable grounds to believe that the Company will be able to pay its debts as and when they become due and payable.

This declaration has been made after receiving the declarations required to be made to the Directors in accordance with Section 295A of the Corporations Act 2001 for the financial year ended 30 June 2015.

Mr Robert W Moses

Independent Non-Executive Chairman

Dated: This-the 21st Day of August 2015.

Mr Mark Diamond

Managing Director



Independent Auditor's Report



Ernst & Young 8 Exhibition Street Melbourne VIC 3000 Australia GPO Box 67 Melbourne VIC 3001 Tel: +61 3 9288 8000 Fax: +61 3 8650 7777 ey.com/au

Independent auditor's report to the members of Antisense Therapeutics Limited

Report on the financial report

We have audited the accompanying financial report of Antisense Therapeutics Limited, which comprises the consolidated statement of financial position as at 30 June 2015, the consolidated income statement and consolidated statement of comprehensive income, the consolidated statement of changes in equity and the consolidated statement of cash flows for the year then ended, notes comprising a summary of significant accounting policies and other explanatory information, and the directors' declaration of the consolidated entity comprising the company and the entities it controlled at the year's end or from time to time during the financial year.

Directors' responsibility for the financial report

The directors of the company are responsible for the preparation of the financial report that gives a true and fair view in accordance with Australian Accounting Standards and the Corporations Act 2001 and for such internal controls as the directors determine are necessary to enable the preparation of the financial report that is free from material misstatement, whether due to fraud or error. In Note 1, the directors also state, in accordance with Accounting Standard AASB 101 Presentation of Financial Statements, that the financial statements comply with International Financial Reporting Standards.

Auditor's responsibility

Our responsibility is to express an opinion on the financial report based on our audit. We conducted our audit in accordance with Australian Auditing Standards. Those standards require that we comply with relevant ethical requirements relating to audit engagements and plan and perform the audit to obtain reasonable assurance about whether the financial report is free from material misstatement.

An audit involves performing procedures to obtain audit evidence about the amounts and disclosures in the financial report. The procedures selected depend on the auditor's judgment, including the assessment of the risks of material misstatement of the financial report, whether due to fraud or error. In making those risk assessments, the auditor considers internal controls relevant to the entity's preparation and fair presentation of the financial report in order to design audit procedures that are appropriate in the circumstances, but not for the purpose of expressing an opinion on the effectiveness of the entity's internal controls. An audit also includes evaluating the appropriateness of accounting policies used and the reasonableness of accounting estimates made by the directors, as well as evaluating the overall presentation of the financial report.

We believe that the audit evidence we have obtained is sufficient and appropriate to provide a basis for our audit opinion.

Independence

In conducting our audit we have complied with the independence requirements of the Corporations Act 2001. We have given to the directors of the company a written Auditor's Independence Declaration, a copy of which is included in the directors' report.

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Opinion

In our opinion:

- a. the financial report of Antisense Therapeutics Limited is in accordance with the Corporations Act 2001, including:
 - giving a true and fair view of the consolidated entity's financial position as at 30 June 2015 and of its performance for the year ended on that date; and
 - complying with Australian Accounting Standards and the Corporations Regulations 2001;
- the financial report also complies with International Financial Reporting Standards as disclosed in Note 1.

Report on the remuneration report

Frank & Young

We have audited the Remuneration Report included in the directors' report for the year ended 30 June 2015. The directors of the company are responsible for the preparation and presentation of the Remuneration Report in accordance with section 300A of the Corporations Act 2001. Our responsibility is to express an opinion on the Remuneration Report, based on our audit conducted in accordance with Australian Auditing Standards.

Opinion

In our opinion, the Remuneration Report of Antisense Therapeutics Limited for the year ended 30 June 2015, complies with section 300A of the Corporations Act 2001.

Ernst & Young

Don Brumley Partner Melbourne

21 August 2015

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Company Directory

DIRECTORS

Mr Robert W Moses Mr Mark Diamond Dr Chris Belyea Dr Graham Mitchell Independent Non-Executive Chairman Managing Director Independent Non-Executive Director Independent Non-Executive Director

COMPANY SECRETARY

Mr Phillip Hains

COMPANY

Antisense Therapeutics Limited ABN 41 095 060 745

REGISTERED OFFICE

6-8 Wallace Avenue Toorak, Victoria, 3142

PRINCIPAL PLACE OF BUSINESS

6-8 Wallace Avenue Toorak, Victoria, 3142

Telephone: + 61 (0)3 9827 8999 Fax: + 61 (0)3 9827 1166

SOLICITORS

Minter Ellison Rialto Towers Level 23, 525 Collins Street Melbourne, Victoria, 3000

SHARE REGISTRY

Boardroom Pty Limited Level 12, 225 George Street Sydney, NSW, 2000

Telephone: 1300 737 760

International: +61 (0)2 9290 9600

SECURITIES QUOTED

Australian Securities Exchange

- Ordinary Fully Paid Shares (ASX Code: ANP)

American Depository Receipts (ADR)

Level 1 ADR Program, ADRs are traded in the US

over-the-counter (OTC) market. Ratio: 1 ADR = 20 ordinary shares

Symbol: ATHJY CUSIP: 037183100

WEBSITE

www.antisense.com.au

AUDITORS

Ernst and Young 8 Exhibition Street Melbourne, Victoria, 3000

BANKERS

Commonwealth Bank of Australia Melbourne, Victoria

