• After years of Society campaigning, the Mental Capacity Act finally received parliamentary approval. This legislation will give people with dementia and carers important new legal rights.

• The first edition of our new newsletter written for and by people with dementia was published. We also held the national launch of Alzheimer’s Forum – a website produced by people with dementia.

• We held a parliamentary lobby to protest about charging for care and the National Institute for Health and Clinical Excellence’s (Nice’s) recommendation that the NHS should no longer prescribe drugs for Alzheimer’s disease. This was our biggest lobby to date and it brought the issue to the attention of the public, the national press and MPs.

• We launched the Alzheimer Society’s first nationally accredited training programme designed specifically for care staff working with people with dementia.

• We started raising awareness of vascular dementia, and of how people can reduce their risk of developing it, through a new three-year project called ‘Hearts and Brains’.

• We will establish a brain bank in partnership with the Alzheimer’s Research Trust to support future research into all types of dementia.

• We will use the information we have gathered from people with dementia and carers to persuade Nice to reconsider its guidance on the use of Alzheimer’s drugs.

• We will develop an electronic dementia training programme, bringing quality information and guidance to care workers without the need for expensive training courses and books.

• We will launch an online research journal in order to make research findings widely accessible.

• We will set up a services directorate to ensure that people with dementia and carers receive the same level of care and support throughout England, Wales and Northern Ireland.
I am delighted to introduce this year’s annual review, which I am sure you will agree demonstrates superbly the breadth and extent of our work in our 25th year. The Alzheimer’s Society is of immense importance, providing support, advice and services to thousands of carers and families affected by dementia. We can be proud of what we have achieved in lobbying government to ensure that the mental capacity bill contained provision for people with dementia. Our protestations and active work to overturn the recommendations of the National Institute for Health and Clinical Excellence (Nice) have contributed to a rethink, although we have yet to learn the final outcome. 2004/2005 was a great year – we helped more people, our income grew, public awareness increased and the government was not allowed to forget us!

To ensure that we can remain in the forefront, and for us to continue to grow and provide consistent services and support across England, Wales and Northern Ireland, the Society is undergoing a series of changes. These involve a newer, smaller board of trustees, along with an advisory council where members will be drawn from all parts of the Society and whose purpose will be to assist the board of trustees in the development of policy for its work.

Considerable changes are also underway at the operational level, which will lead to us being fully compliant with best practice. All of this, of course, creates uncertainty for staff and volunteers alike, and I’d like to pay tribute to everyone involved for their support and commitment, including the chief executive and his senior management team, and the volunteers and staff in the branches and regions, without whom the work of the Society would not be possible. Despite personal concerns about changes in the Society, they have continued to provide support and advice to thousands of people with dementia, their families and carers.

I cannot end this brief introduction without mention of three special people who sadly have died this year: Cora Phillips, the Society’s co-founder; Marjorie Stone, former vice chairman; and Baroness Emily Blatch, former vice president. All three symbolised what is so special about the Society – real dedication to the cause combined with energy and integrity. They will all be sadly missed.

Finally, on behalf of all my trustees, a big thank you to all the members, staff and volunteers who have made the Society what it is.

Dr Nicholas Carey FCGI
Chairman
Alzheimer’s Society
Dementia is the major health and social care challenge of the 21st century. It affects over 750,000 people in the UK, including at least 14,000 people from minority ethnic communities.

The number of people with dementia is set to rise steadily as the population ages and as people are diagnosed earlier. It is estimated that by 2010 there will be approximately 840,000 people with dementia in the UK.

Symptoms of dementia include loss of memory, confusion and difficulties with speech and understanding. Dementia is more than just forgetfulness. It is a degenerative condition, which means that the person’s ability to remember, understand, communicate and reason will gradually decline. There are some drugs available that can slow down the symptoms temporarily but there is, as yet, no cure.

The Alzheimer’s Society provides practical support, information and advice for anybody affected by dementia. This includes people with dementia, family carers, friends, relatives and medical and care professionals.

Although just over half of the people diagnosed with dementia have Alzheimer’s disease, there are over 100 different types of dementia, including vascular dementia, dementia with Lewy bodies and fronto-temporal dementia. The Society supports people with any type of dementia. This year, we launched a new project called ‘Hearts and Brains’ to increase awareness and understanding of vascular dementia.

Most people with dementia are over the age of 65, but the Society works hard to reach out to the 18,000 younger people with dementia in the UK. We provide support for people of all ages. Alzheimer’s Awareness Week 2004® focused on the needs of the many children and young people affected by dementia in the family.

Dementia is a global issue, affecting 18 million people worldwide. The Society is a member of Alzheimer’s Disease International and it works closely with dementia charities and organisations in other countries.
Earlier diagnosis, drugs for Alzheimer’s disease and better awareness mean that more and more people with dementia are able and willing to speak out for themselves. Many are telling us – and the government – what they need. They are using their insight and their passion to reach out to others through writing, public speaking, campaigning and building websites.

REACHING OUT TO OTHERS
This year saw the launch of our new newsletter, *Living with dementia*, written by and for people with dementia. Through the newsletter, people with dementia share experiences, tips and messages of support. As soon as the first edition went out, we were inundated with positive feedback and requests for information. ‘It’s like a breath of fresh air,’ wrote one reader. ‘It made me feel more positive.’ Subscriptions already stand at over 2,000 and we are confident that readership will continue to grow.

Another success of the year was the national launch of the Alzheimer’s Forum. Alzheimer’s Forum is a website that a group of people with dementia living in the West Kent area have been writing and maintaining since 2000. It acts as an online meeting place for people with dementia, where they can talk about their experiences, share a joke and pass on information. This year the group decided that, in order to reach as many people as possible, the forum should be launched on our national website. Please visit www.alzheimers.org.uk/alzheimersforum and take a look.

Much of the Society’s campaigning success is due to the passion and the courage of the people with dementia who support our aims. Throughout the year, people have been writing letters to their MPs to demand better services and access to drugs, speaking publicly about their experiences and taking part in protests and lobbies. This is a powerful way to raise awareness, and increasing numbers of people are keen to speak out.

We held a training session in Kent this year to encourage people who want to work with the media and to give them some experience of how it feels. One brave volunteer agreed to take part in a videotaped interview with a member of our public affairs team. Afterwards he commented, ‘You quickly forget you have an audience and a camera pointing in your face. With an understanding interviewer, you just focus on the conversation and your story.’

People with dementia have also helped us to develop a comprehensive range of helpful and accessible information for anyone experiencing memory problems. *I’m told I have dementia* is a practical guide produced by and for people with dementia. A range of 16 information sheets accompanies the booklet, covering topics such as treatments for dementia, staying healthy and driving.

Across England, Wales and Northern Ireland our 265 Society branches and support groups work directly with people with dementia in the local area, providing services that make dementia easier to live with. Thanks to a generous donation from the Big Lottery Fund, we have been able to employ two development workers for our Living with Dementia programme. They will provide advice and guidance to local branches to enable them to respond well when people with dementia contact them asking for information or support or wanting to get involved in the Society’s work. Some branches run day care or home care services, others run counselling sessions for people with dementia, or organise befriending or advocacy programmes. Many branches run social get-togethers where people with dementia and their carers can enjoy some quality time together.

The North Somerset branch has been holding regular tea dances for its members for the last three years. There is a mix of dance and sing-alongs, followed by tea and sandwiches. Roy Hopkins, who has dementia, comes with his wife Winifred. He says, ‘I was pretty nervous when I came to the tea dance the first time because I’d never really danced much before but I’ve enjoyed coming with Winifred, who likes waltzes.’
Many people are surprised to discover that dementia does not only affect people in their old age. Over 18,000 people with dementia in the UK are under the age of 65. The challenges of dementia are different for younger people – they may have jobs or dependent children and be more physically active. We can provide information about the types of dementia that affect younger people and the services and support available around the country. This year we published Younger people with dementia: an approach for the future, which discusses the needs of younger people and how organisations can meet these needs by working together to provide positive services. The ACE club in Rhyl, funded by the Society’s Rhyl and District branch and the Welsh Assembly’s Carers Strategy, organises a range of activities for younger people. These have ranged from a belly dancing class and Reiki healing to shopping trips to Chester and a ferry across the Mersey. Linda, who attends the club with her husband Gary, says, ‘It’s made a huge difference to me. Before, I was afraid of going outside the house. I was shocked to discover I had dementia because I thought it only affected older people.’

Alzheimer’s disease is not the only cause of dementia. There are over 100 different types of dementia, and vascular dementia is the second most common of these. This year, we launched a three-year project called Hearts and Brains to raise awareness of vascular dementia, especially among people at risk of developing the condition. Vascular dementia is the second most common form of dementia. People with vascular dementia have shared their experiences and worked with us to produce a range of publications that explain the symptoms, progression and treatment of the condition. We will be holding workshops for health professionals all round the country in the coming year to review vascular dementia, the risk factors and the potential for prevention.

OUR NEW LIVING WITH DEMENTIA NEWSLETTER HAD 2,000 SUBSCRIPTIONS WITHIN TWO MONTHS OF BEING LAUNCHED
'Twenty-five years ago my mother went to live in a nursing home. She was just 57 years old. Two years earlier, she had been diagnosed by her GP with ‘pre-senile dementia’. The GP told us it would get worse and that there was no cure. We were devastated and confused. No help or advice was offered and we felt very alone.

Then we saw a programme on television which told the story of a lady who seemed to have the same condition as Mum. They called it Alzheimer’s disease – it was the first time we had heard the term. The programme gave details of a support group that had recently been formed. And so began my long association with the Society.

My beloved mum died in 1988, when the term ‘Alzheimer’s disease’ was still relatively unknown. Things are different now – but I want to thank the Society for being there 25 years ago when there was no one else who understood our pain.'
Hundreds of thousands of people in the UK work tirelessly, caring for friends or relatives who have dementia. It can be an exhausting and demanding task, for which carers often receive little recognition, support or financial help. Nobody should face dementia alone. We are committed to providing carers with practical assistance, emotional support and the information they need to make informed choices.
Many carers feel lonely or isolated at times. It can be difficult to talk about the pressures and anxieties of caring with people who don’t really understand. On our Alzheimer’s Helpline, carers can speak with trained advisers who can provide reassurance and a sympathetic listening ear, as well as give up-to-date information on all aspects of dementia and caring. Last year, the helpline service dealt with around 20,000 enquiries.

Our Lesbian and Gay Network continued throughout the year to provide telephone support for lesbians and gay men caring for a partner, relative or friend. Lesbian and gay carers can face ignorance and prejudice when they try to get access to support and services, adding pressure to an already difficult situation. Knowing that somebody is only a phone call away really can make all the difference.

For more information about the Lesbian and Gay Network please visit www.alzheimers.org.uk/Gay_carers.

We produce a wide range of helpful information sheets, leaflets and booklets. Our monthly newsletter, Share, keeps members informed about dementia, caring, news and research developments. Information is available in large print, in Braille and on audiocassette, and basic information is available in a variety of languages.

Our website contains a wealth of information on dementia, caring, research, news and campaigns. Last year, it received almost one million visits. Please go to www.alzheimers.org.uk to find out more. The website also hosts Talking Point, our online discussion forum. It has almost 1,000 members and is an ideal place to ask for advice, share information, join discussions or simply let off steam! Norman, who cares for his wife Peggy, says, ‘Talking Point (TP) is a rock, an escape, a little world of friends removed from the cares of the day, something to hold on to and look forward to at the end of each day. TP is there for us to rant, to sympathise, to open our hearts and in return find some comfort with our big family. There is always someone there for us.’ To join the TP family please visit www.alzheimers.org.uk/Talkingpoint.

We also provide practical, hands-on assistance for carers all around the country, including befriending projects, training courses and respite care. All Society branches run carers support services; there are about 600 Society carers support groups in the UK. These act as a lifeline for many carers, enabling them to enjoy a change of scene, some company, a chance to talk through their problems, or simply some time to have a cup of tea and put their feet up.

For example, the Bradford branch has run a support group for carers of younger people with dementia for the past five years. The unusual thing about this group is that it meets in a local pub, creating a relaxed and welcoming atmosphere. Carers can bring their loved ones with them – they all enjoy some time together and then split into two separate groups part-way through the evening. Dave, whose wife Maureen has dementia, says, ‘What’s nice about this group is it gives me the chance to chat with others who are in a similar situation and, because it’s a pub, it’s a relaxed setting.’

Many carers struggle to make ends meet. Our dementia support grants scheme aims to relieve some of the immediate financial pressure and to provide encouragement to continue when times are tough. As one carer wrote to us, ‘When my husband became ill, it seemed like a final blow to have financial worries.’ We were able to help this carer with a grant towards the cost of her household bills. She told us later: ‘We are so grateful for your contribution. This certainly supports my determination to keep my darling husband at home for his entire life.’ This year we extended the scheme to make grants available to carers whose loved ones have died or gone into residential care.

THE ALZHEIMER’S HELPLINE DEALS WITH 20,000 ENQUIRIES EVERY YEAR

‘After my phone call to you I felt helped, comforted and reassured’
WE ARE COMMITTED TO SPENDING OVER £1 MILLION ON RESEARCH EVERY YEAR

IN SEARCH OF THE ANSWER

Quality Research in Dementia (QRD) is the award-winning research grants programme of the Alzheimer’s Society. Uniquely, it truly represents the priorities of people with dementia and carers by actively involving them in all aspects of its work.
This mutually beneficial partnership between people with dementia, carers and the research community is made possible through the QRD Network, which lies at the heart of our success. It is made up of 150 people with dementia, carers and former carers. They set the strategy for research, comment on grant applications, select applications for funding, monitor projects and tell people about research results. Their personal experience of dementia means that they know better than anyone which research developments could make the most difference.

With the generous donations of our supporters, we are able to invest over £1 million in research every year. QRD funds research in three different areas: cause, cure and care. The search for cause and cure gives us hope for the future, while research into care ensures that we deliver the best help right now. Research priorities include stem cell research, early diagnosis, risk factors, vaccines, and alternative and complementary therapies. However, we receive many more grant applications than we are able to fund. The success of our grants programme depends on the generosity of donors and supporters.

As Clive Ballard, director of research, says, ‘We should be optimistic that research can deliver improvements in treatment and care in the near future. This can, however, only be achieved if the research strategy is coherent, if the funding is adequate and if there are enough scientists with the right skills.’

For more information, visit www.qrd.alzheimers.org.uk.

We have been able to fund some exciting research projects in the last year, any one of which could lead to dramatic improvements in the quality of life of thousands of people. For example, a generous donation from the Mike Gooley Trailfinders Charity has enabled us to award a three-year fellowship grant to Dr Wendy Noble, who will be investigating why brain cells die in Alzheimer’s disease. She will be looking at how plaques and tangles develop in the brain cells of people with Alzheimer’s disease and how this process could be prevented. This research has already led to a clinical trial of lithium for the treatment of Alzheimer’s disease and is likely to inform the development of other effective new drugs.

It is important that we build and maintain a high profile for dementia research and that dementia scientists are encouraged to share their knowledge and learn from each other. Five of our research fellows were fortunate enough to attend the ninth international conference for Alzheimer’s and related diseases, held in Philadelphia, USA, this year. This was a prestigious event, with 4,500 researchers and old age psychiatrists presenting their work, running sessions and giving talks. One of our research fellows said: ‘We met many researchers, including some who showed interest in future collaboration. By the end of the conference I had a notebook filled with comments, ideas and tutorial notes.’ Disseminating information is a key part of building a high profile for dementia research. We are currently developing an online journal, which will include a variety of review articles on key areas of research. The journal will be freely available and will be launched in the autumn of 2005.

Another key priority for the coming year is the development of an Alzheimer’s brain bank, which will support future work into all types of dementia. We are delighted to be able to work with the Alzheimer’s Research Trust on this exciting project.

‘I cared for my husband for eight years. The Alzheimer’s Helpline helped me so much then that I really wanted to give something back. QRD was a perfect way for me to get involved as I could do the work from home. Over time, I got more and more involved. I am now regional co-ordinator for Wales and I have also been involved in award panels and monitoring groups. The work is varied and really interesting.

Dementia research is so important. I feel very strongly that we should be promoting it as much as possible. We need more effective treatments, and hopefully one day we’ll even find a cure. Then people won’t have to suffer as my husband and I have.’ (Victoria Morgan)
In March 2005, 600 supporters of the Alzheimer’s Society gathered in London for the biggest parliamentary lobby we have ever organised, under the banner of ‘Remember those who forget’. Many were angry and disappointed – but they were also passionate and determined to be heard.

The reason for this outburst of anger and passion? The National Institute for Health and Clinical Excellence (Nice) had just made its shocking recommendation that the NHS should no longer prescribe drugs for Alzheimer’s disease – its argument being that £2.50 a day is too expensive. We have campaigned for many years for the widespread availability of these life-changing drugs. It is unthinkable that Nice should now recommend their withdrawal. Thankfully, we have thousands of vocal and committed supporters, who have deluged Nice with their passionate protests. As actor Tony Robinson says, ‘From my own personal experience I know that dementia doesn’t discriminate – so why should the NHS? I am astonished that Nice thinks that £2.50 a day is too much to pay for precious extra quality time with a loved one with dementia.’

The lobby really pressed home the point and we will not ease the pressure until we have persuaded Nice to reconsider its decision.

The parliamentary lobby also had a second important goal: to call for an end to the charging of personal care. Six years after the Royal Commission on Long Term Care recommended that people with dementia should receive all their care free of charge, people are still expected to pay for help with eating, dressing and using the toilet. Barbara Pointon, carer and speaker at the lobby, says, ‘No way is it ‘personal/social care’ only; attending to hygiene, food and toileting is just the tip of the iceberg. What about the 24-hour vigilance, dealing with changes in personality, difficulties in communication, having the patience to reply to repetitive questions?’ The sheer size of the crowd calling for an end to this injustice said it all, and many people with dementia and carers shared their personal experiences with MPs. The day was a huge success, and we are hopeful that it will bring about much-needed change.

Years of hard work and campaigning paid off in March 2005 when MPs voted the mental capacity bill through its final parliamentary stages. The new Mental Capacity Act marks a major turning point in the legal rights of people with dementia and carers. For the first time, people with dementia cannot legally be presumed incapable of making their own decisions. Carers, too, will benefit from new rights under the act: implementation of the act in 2007 will mean that they must be consulted about decisions relating to the health and welfare of people who have lost capacity. We are delighted that the act has finally become law.

For more information about our campaigning work please visit www.alzheimers.org.uk/News_and_Campaigns. Alzheimer’s Awareness Week 2004® focused on children and young people. When someone is diagnosed with dementia, their children or grandchildren may find it difficult to understand the condition and deal with painful changes in their relationship with loved family members. One young person told us, ‘When my grandfather forgot my name I felt like he didn’t actually love me anymore. But then I found out it was because of dementia.’ We produced an engaging leaflet to explain dementia to children, and sent branches all around the country packs of materials to use in local schools. We also made a video called ‘About my grandfather... about my grandmother’, which premiered in Leicester Square in July. And to top it all, we designed a drawing competition for children, which asked them to show how they thought memories, feelings and thoughts were stored in the brain. The results were colourful, fascinating and often moving.
Thankfully, there are drugs which can ease the symptoms – for a while. Now it seems that small light is about to be extinguished for so many families. £2.50 isn’t much to change someone’s life for a day, is it?”

(Fiona Phillips)
Peter’s story

“I am 80 years of age. I have multiple infarct/vascular dementia, and I know I won’t get better.

I served in the Royal Navy during the Second World War. We used to escort vessels – we did the run from Aden to Bombay and Karachi. I have eight campaign medals and I also have an MBE for my work in the community – I was secretary of the local residents association for many years and I worked with the local police force.

Today, with my health getting worse, I say it’s payback time. The wartime generation, which helped to protect this country, is being betrayed by the government’s policy on charging for care. That’s why I wrote to my MP – if nobody does anything, the government will think it’s alright. People often moan and groan and say to me, “What are we going to do about it?” I say, “Well, what are you going to do about it?”
We are proud of our unique understanding of the needs of people with dementia and carers. We have developed our expertise over many years of research and experience. But this knowledge only becomes truly powerful when it is shared with the people and organisations that can really make a difference.
We share our knowledge through a comprehensive training programme for both Society staff and external organisations. We also produce books, training manuals and videos, and organise conferences.

This year, we launched the Alzheimer’s Society’s first ever nationally accredited training course aimed specifically at improving the quality of care given to people with dementia. The course, ‘An introduction to dementia care’, was produced in collaboration with Bupa Care Homes, Joseph Rowntree Foundation and the University of Sunderland. It uses video sessions and an interactive course workbook to encourage care workers to develop a person-centred and positive approach towards their work. We have high hopes that this course will encourage thousands of care staff to learn more about dementia and take pride in the vital job they do. Graham Stokes, head of mental health at Bupa Care Homes, says, ‘The aim of this programme is to provide carers with an understanding of what it is like to live with dementia and encourage them to view dementia care as a career choice.’

There also continues to be a great demand for our more established training courses, and organisations are increasingly asking us to provide courses for them in-house. This year, Brunelcare—an organisation that provides home care, day care and nursing residential care throughout Bristol and Avon—asked us to provide our six-day ‘Dementia care training skills’ course for its employees. Christine Allen from Brunelcare says, ‘The course was even better than I expected, and in a style I could really relate to.’

We can also put care providers in touch with leading dementia care trainers through our approved trainers scheme. All Society approved trainers have been through a rigorous assessment process and are well qualified to transform an organisation’s training programme and significantly improve the quality of dementia care provision.

It is important that those at the frontline of care – care assistants as well as managers – have easy and direct access to resources to develop their knowledge and skills. Next year we plan to produce a booklet to introduce new care workers to the rewards and the challenges of their chosen career. We also intend to develop an interactive online training programme for both students and care staff, so that they can learn on their own initiative, without the need for expensive training courses and books. Motivating and inspiring those who provide care for people with dementia—and those who will do so in coming years—is a key part of what we do. We are committed to finding the funds and the resources to make this possible.

For more information about the Quality Care team, please visit www.alzheimers.org.uk/Working_with_people_with_dementia.

Publications
Our publications are widely used in a variety of care settings to provide care staff with practical information and guidance on all aspects of dementia care.

- Quality dementia care in care homes: person-centred standards
- Home care services for people with dementia: quality standards
- The Alzheimer’s Society book of activities
- Making each day count: a guide to day care services for people with dementia
- Make a difference in dementia care training: resources that are effective, meaningful and fun!
- Yesterday, today, tomorrow, a video training programme for care staff
- Building on strengths: providing support, care planning and risk assessment for people with dementia
- Food for thought: acute care/care homes/domiciliary care/day care practice guides

This year, almost 2,500 people successfully completed our Yesterday, today, tomorrow certification training scheme.
Our work is dependent on the generosity of our donors and funders and on the commitment of Society supporters all around the country who work so hard to raise money for us. Their continuing support brings us closer to achieving our goals: to support every person with dementia and carer who needs our help, and to find a cure.

RAISING VITAL FUNDS

£30 BUYS OVER TWO HOURS OF A RESEARCHER’S TIME

Donations are the lifeblood of the Society. Every single donation makes a difference and is gratefully received. Many people donate to their local branch in memory of a loved one, or remember us in their will; others give us a one-off cash gift or support us regularly through payroll giving, membership donations or standing orders. To find out about making a donation or becoming a member, please visit www.alzheimers.org.uk/Supporting_the_Society or call the donations hotline on 0845 306 0898.

Local fundraising enables branches to run services for people with dementia and carers. Our nationwide programme of Memory Walks, held to coincide with World Alzheimer’s Day, has become a popular and successful way for branches to raise money. Last year, 4,500 people from 72 branches around England, Wales and Northern Ireland stepped out in support of the Society, raising a total of £134,000. The London Memory Walk took 250 people along a riverside route past some of the country’s best-known landmarks, including Big Ben and the London Eye. The writer and critic AA Gill, who joined the walk, said, ‘Alzheimer’s is a cruel disease... Everyone who has taken part in this year’s Memory Walk has helped raise funds that will make an enormous difference to the lives of people with dementia and their carers.’

Every year, many of our more adventurous supporters complete mountain hikes, run marathons and throw themselves out of aeroplanes to raise money for us. Society supporter David Grigson ran the Great North Run last year and raised a staggering £20,173 for the Alzheimer’s Society – the most ever raised by a single participant in any event. David says, ‘My father had Alzheimer’s disease and after seeing how hard it was for my mother to cope I really wanted to do something for the Society.’ He later ran the London Marathon, raising a further £14,000!

This year, we launched a series of overseas treks to raise money for the...
Society. In spring 2005, 52 explorers set off to trek the Inca Trail in Peru. Everyone completed the trek and the group raised over £150,000. One trek participant said, ‘What a truly memorable trip, I will never forget the experience. Thanks for the opportunity not only to have the journey of a lifetime but also to raise an incredible amount of money for a charity so close to my heart.’ Another 90 people are now eagerly anticipating trips to China in September and October.

Many companies give us valuable income and help to raise awareness of dementia around the country. This year, we have been delighted to work with the Foresters, who have raised £90,000 for us over the last three years, and HBOS, who ran a pin badge campaign in July. Barlow, Lyde & Gilbert also supported us for a second year and organised a fantastic quiz night for us, hosted by Jo Brand. We are extremely grateful to all the wonderful companies who continue to support our work.

We are also grateful to the hundreds of charitable trusts and foundations for their generous donations. The Ingram Trust, for example, has supported the Society for the past six years. It is currently helping to fund our caring grants scheme and our Living with Dementia programme – two crucial areas of our work.

We were also thrilled this year to receive a charitable donation of more than £1 million from the Mike Gooley Trailfinders Charity. The money will be used over the next three years to fund two research fellowships that will improve understanding of dementia and explore ways of improving drug treatments for the condition. Professor Clive Ballard, director of research, said, ‘We are delighted to accept this generous donation. It gives us a fantastic opportunity to build on the major advances that have been made in dementia research since the launch of the Society 25 years ago.’
Society money
In 2005, the Society’s total income from its national organisation, branches and subsidiaries increased by 9 per cent to £33.3 million.

Subscriptions, donations and fundraising income, which accounts for 40 per cent of total income, grew by over £2 million. Over £1 million of this increase was raised in branches with the help of the Society’s branch volunteers and staff.

The Society received restricted funds of £400,000 during the year as part of a commitment of over £1 million from the Mike Gooley Trailfinders Charity.

The launch during the year of overseas treks and the continued success of the Society’s raffle contributed to an increase in surpluses in Alzheimer’s Trading Ltd of more than 50 per cent when compared to the previous year.

Investment income and profits also increased by 46 per cent.

All these successes enabled the Society to spend an extra £2.1 million on its charitable activities during 2004/2005.

Challenges
Even with over 260 branches and support groups, there is more to do. The Society is always looking for ways to help people with dementia and their carers from the moment they need support.

The Society uses its money to top up the shortfalls experienced by many of its contracted services – but it continues to challenge the government commitment for a full cost recovery in future.

The Society is undergoing a series of changes, as part of its One Society programme. The major phases of the implementation of this programme will take place over the coming 18 months. The programme aims to provide more and better services for people with dementia wherever they live in England, Wales and Northern Ireland. It is also an investment in the Society’s infrastructure to ensure that it can continue to improve quality of life for all those touched by dementia well into the future.

As a volunteer-dependent organisation, the Society strives to support and retain volunteers to achieve its objectives. Ever increasing legislation puts considerable pressure on the Society’s resources – it could achieve so much more if this was relieved. As the One Society programme is implemented, it should release volunteers to focus on what they do best.
Reporting on summarised financial statements

The information on this page is extracted from the full trustees’ report and financial statements which have been audited by Horwarth Clark Whitehill LLP who gave an unqualified audit report on 22 August 2005. The auditors have confirmed to the trustees that the summarised financial statements are consistent with the full financial statements for the year ended 31 March 2005. The trustees’ report and financial statements were approved by the trustees and signed on their behalf on 22 August 2005. They will subsequently be submitted to the Charity Commission and the Registrar of Companies.

These summarised financial statements may not contain sufficient information to gain a complete understanding of the financial affairs of the charity. The full trustees’ report, audit report and financial statements may be obtained from the company secretary at Gordon House, 10 Greencoat Place, London SW1P 1PH.

Signed on behalf of the trustees

NAD Carey
Chairman

Graham Dewhirst
Honorary Treasurer

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<td>Membership contributions</td>
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<td>Investment income</td>
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<td>Other income (including trading)</td>
<td>657</td>
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<td><strong>Total unrestricted income</strong></td>
<td>18,360</td>
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<td><strong>RESTRICTED INCOME</strong></td>
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<td>Grants and contracts</td>
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<td>Other income</td>
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<td>Information and education</td>
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<td>Fundraising</td>
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<td><strong>Total expenditure</strong></td>
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<td>Creditors – amounts falling due within one year</td>
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<td>(2,928)</td>
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<td>Creditors – amounts falling due after one year</td>
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<td><strong>Net assets</strong></td>
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<tr>
<td><strong>Total reserves to support expenditure</strong></td>
<td>22,175</td>
<td>19,003</td>
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</table>
Individuals
Ms Lynda Bellingham
Mr Nicholas Bence-Trower
Mr Paul Brooks
Sir Ronald and Lady Cohen
Dame Judi Dench
Ms Nicola Duffett
Ms Britt Ekland
Mr Brian Fisher
Mr AA Gill
Mr Russell Grant
Mr David Grigson
Mr John Hansford
Mr P Hitchman
Mr C Hunt
Ms Sally Lindsay
Mr Phillip McIver
Dr and Mrs JA McShane
Lord Moser
Mr WN Mullings
Mr MHL Paisley
Lady Panufnik
Mrs Joy Passman
Mr John Penrose
Ms Fiona Phillips
Ms Claire Rayner
Mr RJ Rees
Mr Bill Reinking
Ms Anne Robinson
Mr Tony Robinson
Mr and Mrs R Scott
Mr P Tacon
Mr and Mrs D Walmsley
Ms Nicola Wheeler
Mrs Joan Woodward

Associations, groups and clubs
Challenge Adventure Charities
The Marches Highland Ball

Companies
AC Nielsen
AstraZeneca plc
Barlow Lyde & Gilbert
Club La Costa Management Services
Eisai Ltd
Foresters
GMS Estates Limited
Go Ahead Group
HBOS PLC
Laing O’Rourke
Lundbeck Ltd
Mercer
New Covent Garden Food Co
NM Rothschild & Sons
Novartis Ltd
Pfizer Ltd
Powersource Projects
Samsung Electronics
Shire Ltd
Star Cargo plc
Unique Pub Company
Yule Catto & Co plc

Trusts and foundations
29th May 1961 Charitable Trust
Accenture Foundation
Alan James Henry Memorial Trust
Albert Hunt Trust
Alexander Moncur Trust
Amelia Chadwick Trust
Astor Foundation
Beatrice Laing Trust
Bernard Piggott Trust
Bill Brown’s Charitable Settlement of 1989
Blyth Watson Charitable Trust
Bothwell Charitable Trust
Buckland Charitable Trust
CHK Charities Limited
Charles and Amelia Fuchs Memorial Trust
Charles Skey Charitable Trust
Charles Wolfson Charitable Trust
Comic Relief
Constance Travis Charitable Trust
Cowley Charitable Foundation
D & M McCallum Trust
Dame Thora Hird Charitable Trust
Daneway Charitable Trust
David Brooke Charity
Dischma Charitable Trust
Dunhill Medical Trust
Edgar E Lawley Foundation
Elizabeth & Prince Zaiger Trust
Epigioni Trust
Esme Fairbairn Charitable Trust
Excel Fund
FAF Charitable Trust
FG Woodger Trust
Florian Charitable Trust
Forte Charitable Trust
Furlongs Fund
George Gibson Charitable Trust
Gerald Fogel Charitable Trust
Gordon Gray Trust
Grand Lodge of Mark Master Masons
Gresham Charitable Trust
Gunter Charitable Trust
Hawthorne Charitable Trust
Henry Lumley Charitable Trust
Ian Fleming Charitable Trust
Ingram Trust
Jack and Lily Pasha Charitable Trust
Jane Hodge Foundation
John & Edythe Crosfield Charitable Trust
John Beckwith Charitable Trust
Kate Wilson Oliver Trust
Lady Hind Trust
Little Britain Trust Fund
Lloyds TSB Foundation for England and Wales
Lord and Lady Lurgan Charitable Trust
Lyras Family Charitable Trust
Madeline Mabey Trust
Magnet Charitable Trust
Marie Helen Luen Charitable Trust
Martineau Family Trust
Mary Miskin Charitable Trust
Mike Gooley Trailfinders Charity
Mrs Susanna Peake Charitable Trust
Mrs Waterhouse Charitable Trust
Northern Rock Foundation
Oakdale Trust
Palgrave Brown Foundation
Peacock Charitable Trust
Peter Stebbings Memorial Charity
PF Charitable Trust
Philip & Marjorie Robinson’s Charitable Trust
Pilkington Charitable Trust
RG Gale Charity Trust
Rathbones Trust Company
Red Rose Charitable Trust
Richard Kirkman Charitable Trust
Rita Allen Charitable Trust
Rosetrees Trust
Royal Air Force Benevolent Fund
Scouloudi Foundation
Selig Charitable Trust
ShareGift
Shears Charitable Trust
Simon Whitbread Charitable Trust
Sir Samuel Scott of Yews Trust
Sylvia Aitken Charitable Trust
T & J Meyer Family Foundation
The 140 Trustee Company Limited
The David Saunders Family Charitable Trust
The Herbert & Peter Blagrave Charitable Trust
The Thompson Fund
Thomas J Horne Memorial Trust
Tudor Trust
Twitten Charitable Trust
Valentine Charitable Trust
Wilkinson Charitable Trust
Williamson Benevolent Trust
Willie & Mabel Morris Charitable Trust

Thanks also to many other local authorities and social services departments in England, Wales and Northern Ireland.

We are very grateful for every donation that we receive. However, owing to space limitations, we are unfortunately unable to list all of our donors here.

 Supported by The National Lottery through the Big Lottery Fund
In the introduction to this review our chairman referred to some of the changes that are taking place inside the organisation. These changes are substantial and we need to manage the pace, scale and cost of them carefully. But we do need to see these changes through. Throughout the Society there is recognition that these changes will bring huge benefits.

The benefits will be seen in our capacity to do much more for our cause than we are currently able to do. We will offer new services in places where we currently have no presence. We will be able to put the services that we offer now on a more secure footing and we will be able to ensure quality and performance throughout. We will be able to raise more funds for our work. We will be able to harness the tenacity and determination of our volunteers to press home the vital messages about the policies and services that people with dementia need.

Above all, we will move into a phase where the Alzheimer’s Society is clear about what it is trying to achieve and how it will achieve it. We will be in a position to tell our story more clearly than ever before. We will have an even greater role to play in helping people through the difficult journey of living with dementia.

On behalf of the Society, I would like to extend my best wishes to all who use our services – people with dementia, carers and families, professional – and to our volunteers and staff.

Neil Hunt
Chief Executive
Alzheimer’s Society
### ROYAL PATRON
HRH Princess Alexandra GCVO

### PRESIDENT EMERITUS
Sir Jonathan Miller CBE

### PRESIDENT
Jon Snow

### VICE PRESIDENTS
Katie Boyle  
Angela Browning  
Professor Jim Edwardson  
Dr Nori Graham  
Sir John Hannam  
Dr Anne Hunter OBE  
Morella Kayman  
Dr Christine Kirk  
The Rt Hon Lord Morris PC QSO  
Professor Elaine Murphy  
Anne Robinson  
Professor Martin Rossor  
Dame Rosemary Rue DBE CBE  
Malcolm Wicks MP  
Professor Gordon Wilcock

### CHAIRMAN
Dr Nicholas Carey FCGI

### CHIEF EXECUTIVE
Neil Hunt

### PATRONS
Dame Judi Dench  
Ms Britt Ekland

### Council members who served during the year to 31 March 2005

<table>
<thead>
<tr>
<th>Name</th>
<th>Position</th>
<th>Institution</th>
<th>Location</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dr Joe Blackledge MBE (Midlands)</td>
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<tr>
<td>Beryl Cave (London)</td>
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<tr>
<td>resigned 18 September 2004</td>
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<tr>
<td>Corinne Clarke (North West)</td>
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<tr>
<td>Richard Crace</td>
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<tr>
<td>Peter Downing</td>
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<td>John Finnigan (Trent)</td>
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<td>Allan Graham</td>
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<td>Peter Laycock (Eastern)</td>
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<td>Rosalind Macbeth (South West)</td>
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<td>Bill Megraw (Northern Ireland)</td>
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<tr>
<td>William Mitchell</td>
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<tr>
<td>Marilynne Morgan CB</td>
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<td>Alan Peters</td>
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<tr>
<td>Dr Carolyn Popham (Southern)</td>
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<tr>
<td>elected 18 September 2004</td>
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<tr>
<td>Mervyn Powell (Wales)</td>
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<tr>
<td>Joanne Schofield,</td>
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<td>Judy Thompson (Northern)</td>
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<td>Linda Webber</td>
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<td>John Wood</td>
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<tr>
<td>Dr Nori Graham</td>
<td></td>
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<tr>
<td>Dr Diane Hanger</td>
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<tr>
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<td>Professor Robin Jacoby</td>
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<tr>
<td>Ms Enid Levin</td>
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<tr>
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### MEDICAL AND SCIENTIFIC ADVISORY COMMITTEE

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<th>Name</th>
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<th>Institution</th>
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<tr>
<td>Professor Carol Brayne</td>
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<td>Department of Community Medicine</td>
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<tr>
<td>Professor Alistair Burns (Chairman)</td>
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<tr>
<td>Professor Murna Downs</td>
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<td>Professor in Dementia Studies</td>
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<td>Professor Jim Edwardson</td>
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<tr>
<td>Dr Nick Fox</td>
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<td>Dementia Research Group</td>
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<td>Institute of Neurology</td>
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<tr>
<td>Dr Stephen Gentleman</td>
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<td>Neuroinflammation and Repair Unit</td>
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### HONORARY OFFICERS

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<th>Location</th>
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<tr>
<td>Chairman: Dr Nicholas Carey FCGI</td>
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<tr>
<td>Vice Chairman: Irene Kerr</td>
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<tr>
<td>Treasurer: Graham Dewhirst</td>
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<tr>
<td>Professor Simon Lovestone</td>
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