The study of twin subjects permits the documentation of crude heritability and may promote the identification of specific causal alleles. We believe that at the current time, the chief research advantage of twins as subjects, especially monozygotic twins, is that the commonality of their genetic and cultural identity simplifies the interpretation of biological associations. In order to study genetic and environmental determinants of cancer and chronic diseases, we developed two twin registries, maintained at the University of Southern California: The International Twin Study (ITS) and the California Twin Program (CTP). The ITS is a volunteer registry of twins with cancer and chronic disease consisting of 17,245 twin pairs affected by 17,245 twin pairs affected by cancer and chronic disease, respectively, ascertained by advertising in periodicals from 1980–1991. The CTP is a population-based registry of California-born twin pairs ascertained by linking the California birth records to the State Department of Motor Vehicles. Over 51,000 individual California twins representing 36,965 pairs completed and returned 16-page questionnaires. Cancer diagnoses in the California twins are updated by regular linkage to the California Cancer Registry. Over 5,000 cancer patients are represented in the CTP. Twins from both registries have participated extensively in studies of breast cancer, melanoma, lymphoma, multiple sclerosis, systemic lupus erythematosus, diabetes mellitus type 1, mammographic density, smoking, and other traits and conditions.

Keywords: international twin study, California Twin Program, cancer, chronic disease, etiology, twins

Overview and Mission

Two twin registries were developed at the University of Southern California during the 1980s and 1990s (Cockburn et al., 2001c, 2002, 2006; Mack et al., 2000). One, the International Twin Study (ITS), is a volunteer registry of 17,245 twin pairs affected by cancer or other chronic diseases. The other, the California Twin Program (CTP), is a population-based twin registry of native Californians with information about roughly 75,000 native resident twins. The mission of the registries is to identify twins in order to conduct studies on the etiology of chronic diseases. There is currently no infrastructure support for either registry, and support is maintained via individual hypothesis-based awards. Each registry is further described below.

International Twin Study (ITS)

The ITS is a volunteer-based registry of twin pairs in which at least one member is affected by cancer (12,296 affected pairs) or a chronic disease (4,949 affected pairs), and is created to identify potential subjects for studies of environmental and genetic chronic disease determinants (Table 1) (Mack et al., 2000).
From 1980 to 1991, advertisements were placed in over 300 newspapers and magazines across the United States and Canada, with a circulation of roughly 68 million, seeking ‘twins with cancer’ or ‘twins with chronic conditions.’ There was no age criterion; parents of affected twin children responded on behalf of their children. Direct responses from members of affected twins living within the periodicals’ circulation area made up 79% of responses, from those living outside that area 4%; another 17% of the responding twin pairs were referred by friends or family. Once initial contacts were established, a signed permission to obtain medical records, and for cancer cases, pathology reports and diagnostic slides, was requested. Pathology reports and medical records were reviewed for the majority (71.4%) of twin patients, and histological slides were reviewed and classified for 65% of patients, with higher proportions for specific sites. Follow-up with biannual contact was continued until 1993 for the entire registry and longer for study-specific conditions (multiple sclerosis, breast cancer, hematologic neoplasms, melanoma, systemic lupus erythematosus). Respondents were judged by demographic characteristics to be representative of all North American non-Hispanic white twins with cancer or another chronic disease (Mack et al., 2000). Disease concordance patterns by zygosity could be examined and interpreted with possible selection bias in mind. Heritability has been evaluated for type 1 diabetes mellitus (Kumar et al., 1988, 1993), Hodgkin and non-Hodgkin lymphoma (Mack et al., 1995), childhood cancer (Buckley et al., 1996), systemic lupus erythematosus (Deapen et al., 1992), breast cancer (Mack et al., 2002), and multiple sclerosis (Islam et al. 2006) using this cohort. Approximately 6,000 questionnaires have been collected from pairs with systemic lupus erythematosus, diabetes mellitus, multiple sclerosis, melanoma, and breast, lung, bladder, gastrointestinal, and hematologic neoplasms. Biological samples, including blood, saliva, and tumor specimens, have been collected in the context of disease-specific studies, including those of systemic lupus erythematosus (n = 44 pairs), Hodgkin lymphoma (n = 186 pairs), breast cancer (n = 280 pairs), multiple sclerosis (n = 914 pairs), bladder cancer (n = 126 pairs), and pancreatic cancer (n = 45 surviving monozygotic (MZ) twins). Researchers interested in accessing these twins for collaborations should contact Drs Thomas Mack (tmack@usc.edu) or Ann Hamilton (ahamilt@usc.edu).

### California Twin Program (CTP)

The CTP is a population-based registry of twins born in California between 1908 and 1982 (Cockburn et al., 2001c, 2002, 2006). Twins were identified from California birth records, and contact addresses were obtained by linkage to the records of California Department of Motor Vehicles in 1989, 1998, 1999, 2000, and 2001. During this period there were 256,616 multiple births registered, and 161,109 California residents were identified from linkage to the Department of Motor Vehicles (DMV). Recruitment efforts were carried out in four waves between 1991 and 2000, during which letters and invitation were sent to twins with valid addresses. A 16-page screening questionnaire was sent to members of twin pairs with questions on demographic characteristics, zygosity, growth and development, reproductive history, lifestyle factors, dietary preferences, and medical history. Of 115,733 questionnaires mailed, 51,609 (representing 36,965 pairs) were returned, yielding a response rate of 44.6%, similar to or better than those reported among similarly aged persons in other cohort studies. The following tables (Tables 2 and 3) previously published in this journal (Cockburn et al., 2002) describe the cohorts.

### Zygosity and Sex Distribution of Twins Ascertained by the International Twin Study

<table>
<thead>
<tr>
<th>Zygosity and Sex Distribution of Twins Ascertained by the International Twin Study</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cancer</td>
</tr>
<tr>
<td>--------</td>
</tr>
<tr>
<td>MZ (male)</td>
</tr>
<tr>
<td>MZ (female)</td>
</tr>
<tr>
<td>DZ (male–male)</td>
</tr>
<tr>
<td>DZ (female–female)</td>
</tr>
<tr>
<td>DZ (male–female)</td>
</tr>
<tr>
<td>Unclear sex or zygosity</td>
</tr>
<tr>
<td>Total</td>
</tr>
</tbody>
</table>

Note: 1Cardiovascular disease.

### Response Rates Among Located Individual Twins by Age and Sex, California Twin Program

<table>
<thead>
<tr>
<th>Birth year</th>
<th>Age in 2000</th>
<th>Male</th>
<th>Female</th>
<th>Response rates</th>
</tr>
</thead>
<tbody>
<tr>
<td>1908–1917</td>
<td>83–92</td>
<td>282</td>
<td>56.0%</td>
<td>48</td>
</tr>
<tr>
<td>1918–1927</td>
<td>73–82</td>
<td>1,027</td>
<td>60.4%</td>
<td>267</td>
</tr>
<tr>
<td>1928–1937</td>
<td>63–72</td>
<td>1,668</td>
<td>61.9%</td>
<td>448</td>
</tr>
<tr>
<td>1938–1947</td>
<td>53–62</td>
<td>3,508</td>
<td>53.6%</td>
<td>2,856</td>
</tr>
<tr>
<td>1948–1957</td>
<td>43–52</td>
<td>6,601</td>
<td>44.4%</td>
<td>8,545</td>
</tr>
<tr>
<td>1958–1967</td>
<td>33–42</td>
<td>5,062</td>
<td>28.8%</td>
<td>7,635</td>
</tr>
<tr>
<td>1968–1977</td>
<td>23–32</td>
<td>2,907</td>
<td>19.8%</td>
<td>5,182</td>
</tr>
<tr>
<td>1978–1982</td>
<td>18–22</td>
<td>1,913</td>
<td>15.5%</td>
<td>3,640</td>
</tr>
<tr>
<td>Overall</td>
<td>22,988</td>
<td>32.4%</td>
<td>28,621</td>
<td>34.3%</td>
</tr>
</tbody>
</table>

Note: Reproduced from Cockburn et al. (2002) with editor’s permission.
MZ and dizygotic (DZ) twin pairs showed that both genetic and environmental factors are important determinants (Ursin et al., 2009). We are currently developing proposals to confirm these findings.

Hodgkin Lymphoma

We were the first to show an extremely high risk of young adult Hodgkin lymphoma in co-twins of MZ cases compared with co-twins of DZ cases (Mack et al., 1995). By using the unaffected MZ co-twin of a case as a surrogate, we defined a susceptibility immunophenotype characterized by high inflammatory and low T-helper-1 (Th1) cytokine levels (Cozen et al., 2004b, 2008). We then showed a strong inverse association between early childhood behaviors linked to infectious exposures and risk of young adult Hodgkin lymphoma (Cozen et al., 2009). A genome-wide association scan (GWAS) of young adult Hodgkin lymphoma revealed significant associations with loci in the HLA class II region of the genome (Cozen et al., 2012). A polymorphism in PR domain containing protein 1 (PRDM1) was found to predict second primary breast cancer among Hodgkin lymphoma patients treated with radiation therapy (Best et al., 2011). Studies comparing Epstein–Barr virus load and gut microbiota in Hodgkin lymphoma-discordant pairs are ongoing.

Melanoma

We demonstrated that recall bias could at least partially explain associations between sunbathing and melanoma, especially when the respondent believed that sun exposure was a cause of melanoma (Cockburn et al., 2001a, 2001b). However, associations between ease of sunburning and tanning did not appear to be subject to bias. We then found evidence that mole prevalence in general was largely constant, but the MZ:DZ concordance ratio was also as expected, but the MZ:DZ concordance ratio was also increased by North European ancestry, younger age at diagnosis, and Northern latitude of birth (Islam et al., 2006), indicating gene–environment interaction. We then showed that sun exposure modified the risk of multiple sclerosis within disease-discordant MZ pairs (Islam et al., 2007). The twins were included in a large GWAS of multiple sclerosis that identified numerous risk loci in immune response genes (International Multiple Sclerosis Genetics Consortium et al., 2011). Studies evaluating gene–environment interaction are ongoing.

Multiple Sclerosis

We first demonstrated that twin concordance for multiple sclerosis was higher among MZ compared with DZ twins as expected, but the MZ:DZ concordance ratio was also increased by North European ancestry, younger age at diagnosis, and Northern latitude of birth (Islam et al., 2006), indicating gene–environment interaction. We then showed that sun exposure modified the risk of multiple sclerosis within disease-discordant MZ pairs (Islam et al., 2007). The twins were included in a large GWAS of multiple sclerosis that identified numerous risk loci in immune response genes (International Multiple Sclerosis Genetics Consortium et al., 2011). Studies evaluating gene–environment interaction are ongoing.
Smoking
In a study of smoking initiation and persistence, we found that the strongest influence on initiation was having a twin who smoked, with evidence of modification by closeness (Hamilton et al., 2006). Gender strongly influenced this relationship such that co-twin influence was stronger among female compared with male pairs, presumably on the basis of differential behavior. The only significant determinant of smoking persistence was having a co-twin who continued to smoke. In a second study, we were the first to demonstrate that smoking was directly associated with increased interleukin-5 and interleukin-13 levels (Cozen et al., 2004a).

DNA Methylation
In order to investigate epigenetic mediation of environmental determination of chronic disease in identical twins (Cortessis et al., 2012), we are conducting a study of environmental determinants of DNA methylation in twins and have collected paired blood and saliva samples from each member in over 600 MZ pairs to date. Using an Illumina Infinium platform, we have searched unsuccessfully for differences in the DNA methylation status at 19,000 cytosine loci in whole blood DNA between 25 pairs of MZ twins discordant for multiple sclerosis and between 28 pairs of twins discordant for smoking (unpublished). Preliminary results support differences in DNA methylation patterns by hematopoietic cell subsets separated by flow cytometry prior to DNA extraction.

Other Research
We recently showed a twofold increased risk of infectious mononucleosis in co-twins of MZ compared with DZ cases (Hwang et al., 2012). Twins have participated in studies examining genetic variation in the genes encoding proteins related to catecholamine synthesis and regulation, including tyrosine hydroxylase (Zhang et al., 2004), chromogranin A (Chen et al., 2008), and neuropeptide Y(1) receptor (Wang et al., 2009). Other ongoing projects include a study of the determinants of height differences in MZ pairs, agreement between mothers and twins on childhood experiences, and microbiome studies in twins discordant for traits and disease.

Future Plans
Funding for the resources will be provided by hypothesis-driven projects investigating cancer and chronic disease etiology. We also plan to continue to try to secure infrastructure support through a variety of mechanisms.

Acknowledgments
We acknowledge and thank the twins and their families who take the time to participate in our studies. This work was supported by grants from the National Institutes of Health (R01CA110836, R03CA110836, R01CA58839, P30AG017265), the California Tobacco-Related Disease Research Program (7RT-0134H, 8RT-0107H, 6RT-0354H), and the Leukemia Lymphoma Society (TR6137-07).

References


