Which adolescents develop persistent substance dependence in adulthood? Using populationrepresentative longitudinal data to inform universal risk assessment

M. H. Meier¹*, W. Hall^{2,3}, A. Caspi^{4,5,6}, D. W. Belsky^{7,8}, M. Cerdá⁹, H. L. Harrington^{4,5}, R. Houts^{4,5}, R. Poulton¹⁰ and T. E. Moffitt^{4,5,6}

¹Department of Psychology, Arizona State University, Tempe, AZ, USA

² University of Queensland Centre for Clinical Research, University of Queensland, Brisbane, Australia

³National Addiction Centre, King's College, London, UK

⁴Department of Psychology and Neuroscience, Duke University, Durham, NC, USA

⁵ Department of Psychiatry and Behavioral Sciences, Duke University Medical Center, Durham, NC, USA

⁶Social, Genetic, and Developmental Psychiatry Centre, Institute of Psychiatry, King's College London, UK

⁷ Department of Medicine, Duke University Medical Center, Durham, NC, USA

⁸ Social Science Research Institute, Duke University, Durham, NC, USA

⁹ Department of Emergency Medicine, Violence Prevention Research Program, University of California Davis, Davis, CA, USA

¹⁰Dunedin Multidisciplinary Health and Development Research Unit, Department of Psychology, University of Otago, Dunedin, New Zealand

Background. To our knowledge, there are no universal screening tools for substance dependence that (1) were developed using a population-based sample, (2) estimate total risk briefly and inexpensively by incorporating a relatively small number of well-established risk factors, and (3) aggregate risk factors using a simple algorithm. We created a universal screening tool that incorporates these features to identify adolescents at risk for persistent substance dependence in adulthood.

Method. Participants were members of a representative cohort of 1037 individuals born in Dunedin, New Zealand in 1972–1973 and followed prospectively to age 38 years, with 95% retention. We assessed a small set of childhood and adolescent risk factors: family history of substance dependence, childhood psychopathology (conduct disorder, depression), early exposure to substances, frequent substance use in adolescence, sex, and childhood socioeconomic status. We defined the outcome (persistent substance dependence in adulthood) as dependence on one or more of alcohol, tobacco, cannabis, or hard drugs at \geq 3 assessment ages: 21, 26, 32, and 38 years.

Results. A cumulative risk index, a simple sum of nine childhood and adolescent risk factors, predicted persistent substance dependence in adulthood with considerable accuracy (AUC = 0.80).

Conclusions. A cumulative risk score can accurately predict which adolescents in the general population will develop persistent substance dependence in adulthood.

Received 7 August 2015; Revised 14 October 2015; Accepted 26 October 2015; First published online 1 December 2015

Key words: Longitudinal, risk assessment, substance dependence, universal screen.

Introduction

There is increasing interest in community-based, universal risk assessment to identify youth who either have a substance-use disorder or who will develop one in the future. Universal risk assessment, followed by appropriate intervention, could potentially reduce the population burden of disease associated with substance-use disorders. There are many risk-assessment tools that screen adolescents for current or future substance-use disorder (Clark *et al.* 2006; Vanyukov *et al.* 2009; Chung *et al.* 2012; Kirisci *et al.* 2013; Levy *et al.* 2014). None, to our knowledge, incorporate the key features of the most successful universal risk assessment tools to date, such as the Framingham risk score for cardiovascular disease (Wilson *et al.* 1998; D'Agostino *et al.* 2008). The Framingham risk score was developed using population-based samples, estimates total risk briefly and inexpensively by incorporating a relatively small number of well-established risk factors, and aggregates risk factors using a simple algorithm.

^{*} Address for correspondence: M. H. Meier, PhD., Department of Psychology, Arizona State University, PO Box 871104, Tempe, AZ 85287-1104, USA.

⁽Email: madeline.meier@asu.edu)

In the present study, we developed a risk score to identify adolescents in the general population who are at risk for persistent substance dependence. To maximize translation to practice in community settings, we incorporated the key features of the most successful universal risk assessment tools. That is, we used data from a population-representative longitudinal study; we selected a relatively small number of risk factors that have been shown in longitudinal studies to consistently and robustly predict substance dependence; and we aggregated these risks into a risk score using a simple algorithm, namely the sum of dichotomous risks. This summation approach draws on the large body of research showing that number of childhood risk factors predicts poorer mental and physical health in adulthood (i.e. cumulative risk) (Rutter, 1981; Sameroff et al. 1987; Felitti et al. 1998; Evans et al. 2013).

We evaluated the accuracy with which the cumulative risk score in adolescence predicted risk for substance dependence through young adulthood to early midlife. However, rather than predicting an adolescent's risk for lifetime substance dependence, we predicted risk for severe, persistent substance dependence. Our rationale was that epidemiological studies show that the prevalence of lifetime substance dependence is quite high, and most people with substance dependence remit on their own without treatment (Heyman, 2013; Meier et al. 2013; Grant et al. 2015). Therefore, to avoid over-treating adolescents who would benefit from brief, harm-reduction interventions, and under-treating adolescents who might require more intensive intervention, we developed a population-based risk score that distinguishes those with the poorest long-term prognosis from those with a relatively good prognosis. In addition, rather than predicting risk for dependence on specific substances (e.g. alcohol v. cannabis), we collapsed across substances in defining persistent forms of substance dependence. Our reasoning was that practitioners conducting universal screening (e.g. primary care physicians) want to assess risk for severe dependence on any substance, as opposed to risk for particular types of substance dependence. We further reasoned that the development of substance-specific risk assessment tools would result in the proliferation of risk assessments, thereby reducing implementation in practice.

Method

Participants

Participants are members of the Dunedin Multidisciplinary Health and Development Study, a longitudinal investigation of health and behavior in a complete birth cohort (Poulton et al. 2015). Study members (N = 1037, 91% of eligible births, 52% male) were all individuals born between April 1972 and March 1973 in Dunedin, New Zealand, who were eligible for the longitudinal study based on residence in the province at age 3 and who participated in the first follow-up assessment at age 3. The cohort represents the full range of socioeconomic status (SES) in the general population of New Zealand's South Island and is primarily white (Moffitt et al. 2001). On adult health, the cohort matches the NZ National Health & Nutrition Survey (e.g. body mass index, smoking, general practitioner visits) (Poulton et al. 2006). Assessments occurred at birth and at ages 3, 5, 7, 9, 11, 13, 15, 18, 21, 26, 32, and, most recently, 38 years, when 95% of the living 1007 study members took part. At each assessment phase, study members were brought to the Dunedin Research Unit for a full day of interviews and examinations.

Ethics statement

The study protocol was approved by the institutional ethical review boards of the participating universities. The authors assert that all procedures contributing to this work comply with the ethical standards of the relevant national and institutional committees on human experimentation and with the Helsinki Declaration of 1975, as revised in 2008. The Otago Ethics Committee approved each phase of the study. Informed consent was obtained from all study members.

Persistent substance dependence

Past-year substance dependence diagnoses were made using the Diagnostic Interview Schedule (DIS) following Diagnostic and Statistical Manual of Mental Disorders (DSM) criteria (Robins et al. 1981, 1995; APA, 1987, 1994). We assessed alcohol, tobacco, and cannabis dependence at ages 21, 26, 32, and 38 and hard-drug dependence (e.g. heroin, cocaine) at ages 26, 32, and 38. DSM-III-R criteria were used at age 21 and DSM-IV criteria were used at ages 26, 32, and 38. We have previously compared prevalence rates of alcohol and cannabis dependence in the Dunedin Study with other representative studies of same-age respondents (Moffitt et al. 2010). The past-year prevalence of alcohol dependence was similar in the Dunedin Study, the National Epidemiologic Survey on Alcohol and Related Conditions (NESARC), and the National Comorbidity Survey (NCS; Moffitt et al. 2010). The past-year prevalence of cannabis dependence was slightly higher in the Dunedin Study than in representative U.S. surveys, but was similar to another longitudinal, population-representative survey of New Zealanders (Moffitt et al. 2010). The past-year prevalence of tobacco dependence in the Dunedin

Study (averaged across ages 18–38, 17%) was similar to NESARC participants aged 18+ (13%) (Grant et al. 2004). (NCS did not assess past-year tobacco dependence.) Direct comparison of rates of hard-drug dependence across studies is difficult due to differences between studies in the drugs included in this category. However, in general, rates of hard-drug dependence appear to be slightly higher in the Dunedin Study. To summarize, the prevalence of alcohol and tobacco dependence in the Dunedin Study is similar to other representative U.S. studies, but the prevalence of cannabis and hard-drug dependence is slightly higher in Dunedin. One potential explanation for this is that the prevalence of cannabis and hard-drug dependence is, indeed, higher in New Zealand. Another potential explanation for this pattern of findings is that Dunedin Study participants, interviewed repeatedly over the course of their lives, have learned to trust the study's confidentiality guarantee, and are, therefore, more forthcoming about their illicit drug use.

We classified study members as persistently substance dependent if they were diagnosed with one or more of alcohol, tobacco, cannabis, or hard-drug dependence at 3+ assessment ages (ages 21, 26, 32, 38). For example, a study member could be considered persistently dependent if they were diagnosed with alcohol dependence at 3+ assessment ages (homotypic continuity). A study member could also be considered persistently dependent if they were diagnosed with tobacco dependence at age 21, followed by cannabis dependence at age 26 and hard-drug dependence at age 32 (heterotypic continuity). Of those classified as persistently dependent, 73% both diagnosed persistently for a single substance and across different substances. We chose a threshold of 3+ diagnoses to ensure that we were capturing individuals with severe, chronic dependence throughout adulthood.

We collapsed across substances in defining persistent dependence because practitioners want to predict risk for severe dependence on any substance, rather than dependence on a particular substance. This decision to collapse across substances is bolstered by evidence that (*a*) different substance-use disorders tend to co-occur (Table 1) (Krueger *et al.* 2002; Kendler *et al.* 2003; McGue *et al.* 2006), (*b*) a common liability underlies all substance-use disorders (Krueger *et al.* 2002; Kendler *et al.* 2003; McGue *et al.* 2006) and, (*c*) our results were similar across specific substances (Supplementary Tables S1–S4).

To be classified as persistently substance dependent, study members had to have been assessed for dependence at three of four assessment occasions. Ninety-three percent of the original 1037-member cohort were classified (910 had diagnostic data for four assessment occasions and 961 had data for three). Of those not classified, nearly half (n = 37) had either died or left the study before age 18 or had severe developmental disabilities that prevented their being interviewed with the DIS.

Risk factors

The nine childhood and adolescent risk factors are described in Table 2: SES, family history of substance dependence, conduct disorder, depression, early exposure to substances, frequent alcohol use, frequent tobacco use, frequent cannabis use, and male sex. We selected these particular risk factors because they (i) have been shown in longitudinal studies to consistently and robustly predict adult substance dependence, (ii) represent pre-specified domains of obvious interest (sociodemographic characteristics, mental health, and substance use), and (iii) have fairly natural cut-offs (Grant & Dawson, 1998; Chassin *et al.* 2004; Fergusson *et al.* 2007; Pardini *et al.* 2007; Brook *et al.* 2011; Hussong *et al.* 2011; Stone *et al.* 2012; Kendler *et al.* 2013).

Statistical analysis

We summed the nine dichotomous childhood and adolescent risk factors to produce a single cumulative risk index that allowed us to classify individuals as persistently substance dependent based on their number of risks. We evaluated predictive accuracy using the traditional performance measures: area-under-the-curve, sensitivity, specificity, positive predictive value (PPV), and negative predictive value (NPV). Choosing a higher threshold for number of risks results in lower sensitivity (more false negatives) but higher specificity (fewer false positives). We plotted sensitivity against 1 - specificity for every value of the index, yielding a receiver-operating characteristic (ROC) curve. The area under the curve (AUC) provides a measure of predictive accuracy that reflects the probability of correctly classifying a randomly selected pair of individuals in which one has persistent substance dependence and the other does not. The AUC can take on any value between 0.50 (indicating chance prediction) and 1.00 (indicating perfect prediction). AUC values of 0.54, 0.64, and 0.71 correspond to Cohen's d values of 0.20, 0.50, and 0.80, and reflect small, medium, and large effects, respectively (Rice & Harris, 2005).

Results

The prevalence of persistent adult substance dependence to age 38 in this population-representative cohort was 19% (n = 183). Table 3 shows that each childhood and adolescent risk factor significantly predicted persistent substance dependence in adulthood, and frequent to-bacco use in adolescence was the best predictor (Table 3, left panel; AUC = 0.74). Adolescent tobacco use remained a top predictor even when tobacco

	Study mem	ber category					
Substance dependence	All adults (N=961)	Persistent alcohol dependence (N=40)	Persistent tobacco dependence (N=96)	Persistent cannabis dependence (N=27)	Persistent hard-drug dependence (N = 10)	Persistent substance dependence (N = 183)	Persistent dependence on substances excluding tobacco ^a (N = 83)
Persistent alcohol dependence	4.17	100.00	19.79	25.93	30.00	21.86	48.19
Persistent tobacco dependence	10.02	47.50	100.00	48.15	50.00	53.04	38.55
Persistent cannabis dependence	2.81	17.50	13.54	100.00	30.00	14.75	32.53
Persistent hard-drug dependence	1.07	7.69	5.32	11.54	100.00	5.68	12.35
Lifetime alcohol dependence	31.84	100.00	65.62	70.37	60.00	73.22	85.54
Lifetime tobacco dependence	34.03	72.5	100.00	85.19	70.00	89.07	75.90
Lifetime cannabis dependence	16.34	55.0	42.71	100.00	90.00	53.55	74.70
Lifetime hard-drug dependence	6.56	35.00	21.87	48.15	100.00	26.78	45.78

Table 1. Prevalence of persistent (3+ diagnoses between ages 21 and 38) and lifetime (at least one diagnosis between ages 21 and 38) substance dependence by study member category

This table shows the prevalence of the disorders listed in the rows given the group listed in the columns. For example, of all adults in the cohort (column 1, N = 961), 4.17% had persistent alcohol dependence (row 1). As another example, of those with persistent alcohol dependence (column 2, N = 40), 47.50% had persistent tobacco dependence (row 2).

^a Even after excluding tobacco dependence from the criteria for persistent substance dependence, 38.55% of the group with persistent substance dependence also met criteria for persistent tobacco dependence (last column, second row). This is because many of those who were persistently dependent on alcohol, cannabis, or hard drugs were also persistently dependent on tobacco, and 75.90% (last column, sixth row) had been dependent on tobacco at some point in their life between ages 21 and 38.

Table 2. Childhood and adolescent risk factors

Risk ^a	Respondent	Description ^b	Study member's age(s) at assessment	Brief screen adaption of risk ^c
(1) Low family SES	Parents	The highest of father's or mother's occupation using a 6-point scale for New Zealand (Elley & Irving, 1976). Repeated measures were averaged (Wright <i>et al.</i> 1999). Study members were divided into two groups: high or intermediate SES (manager or physician, secretary or electrician) and low SES (cashier or textile machine operator)	Birth–15	Not included in the brief screen, because this measure is based on data aggregated across 15 years
(2) Family history of substance dependence	Study member and parents	The Family History Screen (Weissman <i>et al.</i> 2000) was used to obtain the proportion of family members, across three generations, with a diagnosis of substance dependence (alcohol or drug dependence) (Milne <i>et al.</i> 2009 <i>a, b</i>). Study members with a proportion of 30% or more were classified as having a family history of substance dependence	32	Maternal reports of alcohol problems for six family members: study members' biological parents and all four grandparents. Mothers answered the following questions: 'Has ever had any treatment or been in hospital for drinking?' 'Has ever had alcoholism?' 'Has ever had a drinking problem or did other people think he/she had a drinking problem?' (Odgers <i>et al.</i> 2007). An affirmative answer to any question was deemed positive for family history
(3) Childhood conduct disorder	Study member	Past-year conduct disorder was assessed using the Diagnostic Interview Schedule for Children (Costello <i>et al.</i> 1982) at ages 11, 13, and 15 and the Diagnostic Interview Schedule (Robins <i>et al.</i> 1981) at age 18. Diagnoses were based on DSM-III criteria at the younger ages and DSM-III-R criteria at age 18. Conduct disorder criteria at each assessment phase were scored to be consistent with DSM-IV (Moffitt <i>et al.</i> 2001)	11, 13, 15, and 18	One symptom was taken from the age 18 conduct disorder assessment. We selected the symptom that correlated most highly with the full diagnosis: breaking in
(4) Childhood depression	Study member	Past-year depression was assessed using the Diagnostic Interview Schedule for Children (Costello <i>et al.</i> 1982) at ages 11, 13, and 15 and the Diagnostic Interview Schedule (Robins <i>et al.</i> 1981) at age 18. Diagnoses were based on DSM-III criteria at the younger ages and DSM-III-R criteria at age 18	11, 13, 15, and 18	One symptom was taken from the age 18 depression assessment. We selected the symptom that correlated most highly with the full diagnosis: fatigue or loss of energy
(5) Early exposure to substances	Study member	Use of drugs (e.g. inhalants, cannabis) or use or purchase of alcohol on multiple occasions over the past year at age 13, age 15, or both (Odgers <i>et al.</i> 2008)	13 and 15	Same
(6) Adolescent frequent alcohol use	Study member	Study members reported on their frequency of alcohol use over the past year at age 18. Study members who reported using alcohol on 5+ days per week were considered frequent alcohol users	18	Same

Risk ^a	Respondent	Description ^b	Study member's age(s) at assessment	Brief screen adaption of risk ^c
(7) Adolescent frequent tobacco	Study member	Study member Study members reported on their frequency of tobacco use over 18 the past year at age 18. Study members who reported using	18	Same
use (8) Adolescent frequent cannabis use	Study member	tobacco on a daily basis were considered frequent tobacco users Study members reported on their frequency of cannabis use over 18 the past year at age 18. Study members who reported using cannabis on 5+ days per week were considered frequent cannabis	18	Same
		users		
SES, Socioeconomic status. ^a Male sex was also include	uic status. so included as a ri	SES, Socioeconomic status. ^a Male sex was also included as a risk factor in the cumulative risk index.		

dependence was excluded from the outcome (Table 3, right panel; AUC = 0.69), which was unsurprising given the high rates of co-morbidity between tobacco dependence and dependence on other substances (Table 1).

The cumulative risk index (mean = 1.78, s.D. = 1.50) predicted persistent substance dependence in adulthood with considerable accuracy. The ROC analysis revealed an AUC of 0.80, a large effect, meaning that we had an 80% probability of correctly predicting, from a randomly selected pair of adolescents, which adolescent would have persistent substance dependence in adulthood. Results were similar when tobacco dependence was excluded from the outcome (Table 3, right panel; AUC = 0.81) and when predicting persistent dependence on each substance individually (Supplementary Tables S1–S4).

The prevalence of persistent substance dependence increased markedly as a function of number of child-hood and adolescent risks (Fig. 1*a*, striped bars): 3% of adolescents with 0 risk factors, 27% of adolescents with 3 risk factors, and 74% of adolescents with 6+ risk factors had persistent substance dependence as adults.

Table 4 shows sensitivity, specificity, PPV, NPV, and overall classification accuracy of the cumulative risk index as a function of number of risks. Overall accuracy was greatest at a cut-off on the cumulative risk index of 4+ to 5+ risks.

Moderation by sex

S6 for a sample brief screen form that could be used in community settings.

^o Measures showed reliability >0.70.

See Supplementary Table

Male sex was included as a risk factor in the cumulative risk index, but to test the possibility that the cumulative risk index is a more accurate predictor of persistent substance dependence for one sex, we examined sex as a moderator. First, we tested whether there were sex differences in the associations between each risk factor in the cumulative risk index and persistent substance dependence. There was no evidence that sex moderated any of these associations. Next, we removed male sex as a risk factor in the cumulative risk index and recomputed the AUC for the cumulative risk index separately by sex. The cumulative risk index (without sex as a risk factor) predicted persistent substance dependence similarly well for girls (AUC= 0.81) and boys (AUC=0.78). Moreover, the cut-off score that maximized overall classification accuracy was 4+ to 5+ risks for girls and 3+ risks for boys, v. 4+ to 5+ risks for both girls and boys when male sex was included as a risk factor in the cumulative risk index. Findings suggest that the cumulative risk index predicts persistent substance dependence similarly for girls and boys. A practical advantage of including male sex as a risk factor in the cumulative risk index is that it equates the cut-off score for girls and boys.

 Table 2 (cont.)

	Persistent dependence on alcohol, tobacco, cannabis, and hard-drugs (19%)				Persistent dependence on substances except tobacco (9%)			
Childhood and adolescent risk factors	RR	95% CI	р	AUC	RR	95% CI	р	AUC
Low family socioeconomic status	1.73	1.32-2.27	< 0.001	0.56	1.80	1.17–2.79	0.008	0.56
Family history of substance dependence	2.62	2.03-3.37	< 0.001	0.61	2.84	1.88-4.27	< 0.001	0.61
Childhood conduct disorder	3.20	2.50-4.09	< 0.001	0.66	5.42	3.58-8.19	< 0.001	0.71
Childhood depression	2.05	1.58-2.67	< 0.001	0.59	2.50	1.65-3.79	< 0.001	0.60
Early exposure to substances	2.91	2.24-3.79	< 0.001	0.60	3.76	2.47-5.73	< 0.001	0.62
Adolescent frequent alcohol use	2.30	1.49-3.54	< 0.001	0.53	2.99	1.56-5.74	0.001	0.54
Adolescent frequent tobacco use	5.41	4.00-7.31	< 0.001	0.74	4.28	2.72-6.74	< 0.001	0.69
Adolescent frequent cannabis use	4.25	3.22-5.61	< 0.001	0.55	9.51	6.49-13.93	< 0.001	0.61
Male	1.54	1.18-2.02	0.002	0.57	2.89	1.79-4.66	< 0.001	0.63
Cumulative risk index				0.80				0.81

Table 3. Relative risk of persistent substance dependence in adulthood given each childhood and adolescent risk factor

RR, Relative risk; CI, confidence interval; AUC, area under the curve.

The risk factors in this table were modestly correlated (*r*'s ranged in absolute magnitude from 0.00 to 0.33). Note that AUC is a better indicator of classification accuracy than relative risk (Pepe *et al.* 2004).

Sensitivity analyses

We tested whether prediction could be improved by adding another risk factor to the cumulative risk index - either low childhood self-control or childhood maltreatment. These risk factors were selected because they have been shown to predict risk for substance dependence and because they were available in the dataset. Both low childhood self-control and childhood maltreatment were associated with increased risk of persistent substance dependence, and they predicted persistent substance dependence with similar accuracy to other predictors (Supplementary Table S5; AUCs = 0.55 and 0.57, respectively). Adding these risk factors to the cumulative risk index did not improve accuracy. When either risk factor was added, the cumulative risk index predicted persistent substance dependence with an AUC of 0.79.

Next we tested the effects of dropping a risk factor from the cumulative risk index. Table 5 shows that the AUC did not drop substantially with the exclusion of any single risk factor, with exception of frequent tobacco use in adolescence.

Finally, we tested the robustness of the cumulative risk index by reducing the threshold for 'persistent' substance dependence from 3+ to 2+ diagnoses across assessment ages. The cumulative risk index was about as accurate when predicting 2+ substance dependence diagnoses from ages 21 to 38 (AUC = 0.77) as when predicting 3+ diagnoses (AUC = 0.80). The cumulative risk index was also fairly accurate when predicting 1+ diagnoses from ages 21 to 38 (AUC = 0.76).

The cut-off score on the cumulative risk index that maximized overall classification accuracy dropped from 4-5+ risks to 3-4+ risks to 2+ risks for predicting 3+, 2+, and 1+ dependence diagnoses, respectively, from ages 21 to 38. This is not surprising given the dose-response association between number of childhood and adolescent risk factors and persistence of substance dependence. For example, the mean number of substance dependence diagnoses from ages 21 to 38 increased in a fairly linear fashion as a function of number of childhood and adolescent risk factors (Fig. 1b). Fig. 1a shows that cohort members with a greater number of risk factors were at higher risk of dependence, regardless of how we defined the dependence outcome (1+, 2+, or 3+ diagnoses). The vast majority of individuals with 0 risks never developed dependence (79%), whereas this was true of only 6% of those with 4+ risks.

Comparison with other risk screens

Both NIDA's Quick Screen and NIAAA's Alcohol Screening Guide for Children and Adolescents rely exclusively (NIDA) or heavily (NIAAA) on assessing frequency of drug use and/or alcohol use. Table 3 and Supplementary Tables S1–S4 show that by expanding risk assessment beyond frequency of drug and alcohol use, prediction was improved. For example, frequent cannabis use in adolescence predicted persistent cannabis dependence in adulthood with an AUC of 0.67, whereas the cumulative risk index predicted persistent cannabis dependence with an AUC of 0.83 (Supplementary

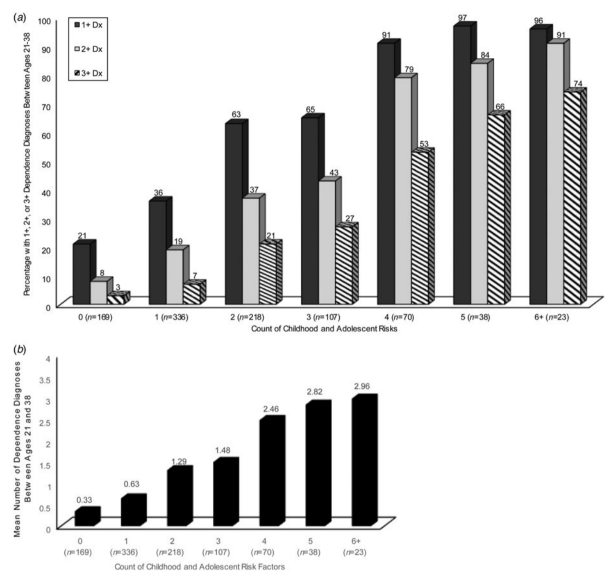


Fig. 1. (*a*) Percentage of the population-representative cohort who had 1+, 2+, or 3+ dependence diagnoses as a function of number of childhood and adolescent risks. Percentages (shown above each bar) can be used to calculate an adolescent's relative risk for persistent adult substance dependence. For example, adolescents with 2 risks were seven times more likely to diagnose with dependence 3+ times between ages 21 and 38 than their peers with 0 risks (21/3=7). (*b*) Mean number of substance dependence diagnoses between ages 21 and 38 as a function of number of childhood and adolescent risks. Note. Number of diagnoses ranged from 0 (was not diagnosed with dependence at age 21, 26, 32, or 38) to 4 (was diagnosed with dependence at age 21, 26, 32, and 38). Mean number of dependence diagnoses is shown above each bar. For example, the mean number of substance dependence diagnoses between ages 21 and 38 for people with 2 risks was 1.29.

Table S3). The cumulative risk index was more accurate in predicting persistent dependence on each substance than frequency of alcohol or drug use alone, with one exception. Frequent tobacco use in adolescence predicted persistent tobacco dependence as well as the cumulative risk index (Supplementary Table S2).

Brief screen adaption

We adapted our risk measures for a brief screen to see whether the cumulative risk index could be used in community settings by practitioners with limited time to conduct the detailed risk assessments in our research study. Table 2 shows how we adapted each measure, and Supplementary Table S6 shows the brief screen. The brief screen (i.e. a cumulative risk index based on the sum of the eight adapted risks) performed nearly as well the cumulative risk index based on our more detailed risk assessments (AUC = 0.79 v. 0.80, respectively). Like the cumulative risk index based on our more detailed risk assessments, the brief screen was more accurate in predicting persistent

No. of risk factors	Sensitivity	Specificity	PPV	NPV	Overall accuracy (%)	Proportion of the populatior with no. of risk factors
1+	0.97	0.21	0.22	0.97	36	82.41
2+	0.84	0.61	0.34	0.94	66	47.45
3+	0.59	0.83	0.45	0.90	79	24.77
4+	0.43	0.93	0.60	0.87	84	13.63
5+	0.23	0.98	0.69	0.84	83	6.35
6+	0.09	0.99	0.74	0.82	82	2.39

Table 4. Sensitivity, specificity, positive predictive value, negative predictive value, and overall prediction accuracy of the cumulative risk index as a function of number of risk factors (n = 961)

PPV, Positive predictive value: the % of those predicted to develop persistent substance dependence (3+ diagnoses of substance dependence between ages 21 and 38) who actually developed persistent substance dependence. NPV, negative predictive value: the % of those predicted not to develop persistent substance dependence who actually did not develop persistent substance dependence. Overall accuracy = the proportion of the sample that was correctly classified. We included study members with missing predictor data in our analysis of the cumulative risk index. Ninety-five percent of the 961 study members had data for 7+ of the nine predictors. When we restricted the analysis to participants with no missing predictor data, results were unchanged.

Table 5. Accuracy of the cumulative risk index in predicting
 persistent substance dependence after systematically removing one

 risk factor from the index
 Image: State of the systematical systema

Risk factor removed from the cumulative risk index	AUC for the cumulative risk index
No risk factors removed	0.80
Low family socioeconomic status	0.80
Family history of substance dependence	0.79
Childhood conduct disorder	0.79
Childhood depression	0.79
Early exposure to substances	0.79
Adolescent frequent alcohol use	0.80
Adolescent frequent tobacco use	0.75
Adolescent frequent cannabis use	0.80
Male	0.78

AUC, Area under the curve.

dependence on each substance than frequency of alcohol or drug use alone with one exception. Frequent tobacco use in adolescence predicted persistent tobacco dependence as well as the brief screen. We report additional information on the accuracy of this brief-screen adapted risk index in Supplementary Table S7 and Supplementary Fig. S1.

Discussion

This report advances knowledge by suggesting answers to three recently posed questions about screening adolescents for risk of substance-use disorders (Subramaniam & Volkow, 2014). The first question is: How can we combine multiple risk factors to

estimate an adolescent's risk? Current screening tools rely heavily on assessing adolescent substance use, yet other risk factors, such as psychiatric disorder, also predict risk (Subramaniam & Volkow, 2014). In this report, we showed that summing a small set of dichotomous risks into a single cumulative risk index is a clinically useful way to integrate risk factors and accurately predict persistent substance dependence. Many studies have shown that cumulative childhood risk is associated with adult mental and physical health problems (Rutter, 1981; Sameroff et al. 1987; Felitti et al. 1998; Evans et al. 2013). A recent study even showed that cumulative risk distinguishes those with persistent alcohol problems from those with time-limited alcohol problems (Copeland et al. 2012). The current report extends this work to suggest how cumulative risk could be used as an actuarial risk assessment tool in community settings to accurately predict persistent substance dependence in the general population.

The second question we address is: Who is at risk? This report provides initial population-representative estimates of risk that can be used to gauge an individual adolescent's likelihood of having persistent substance dependence in adulthood. For example, 3% of adolescents with 0 risks, 27% of adolescents with 3 risks, and 74% of adolescents with 6+ risks developed persistent substance dependence in adulthood. In the future, practitioners may use risk estimates such as these (aggregated across more population-representative studies like ours) to make actuarial judgments and referrals to treatment. One may even envisage members of the general population calculating their risk for persistent substance dependence on their own, just as they now can calculate their risk for heart attack (http://cvdrisk. nhlbi.nih.gov/calculator.asp).

The third question we address is: How should we decide the appropriate level of intervention for an at-risk adolescent? By triaging adolescents into different levels of intervention based on their risk, we might avoid under- or over-treatment and reduce the costs of treatment. The cumulative risk index lends itself to triaging, because we documented a dose-response association between number of childhood and adolescent risk factors and persistence of substance dependence. That is, individuals with more risks had more persistent substance dependence. This dose-response association suggests that adolescents with more risks may require more intensive intervention, whereas individuals with fewer risks may benefit from brief interventions. Although more work will be needed to determine the cut-off scores on the cumulative risk index that decide level of care, we illustrate one possible way that the cumulative risk score could be used.

Adolescents with 4+ risks might be candidates for intensive intervention. A cut-off score of 4+ risks maximized overall accuracy in predicting persistent dependence in adulthood. Moreover, specificity at 4+ risks was high (93%), ensuring that costly interventions go to the small portion of the population (13.6%) that really need it (Table 4). Overtreatment of adolescents with 4+ risks is unlikely, as 94% were diagnosed with dependence at least once between ages 21 and 38 and 82% were diagnosed at least twice. These adolescents may benefit from broad-based interventions (i.e. interventions that target a variety of risk factors, not just substance use) because of the number and variety of their accumulated risks. Interventions need not focus exclusively on the small set of predictors studied here. These predictors were selected for the explicit purpose of efficiently identifying those at highest risk. Intervention design should be guided by the literature on effective treatments that target mutable risks for a variety of problem behaviors.

Brief interventions might be offered to adolescents with 2 and 3 risk factors. These adolescents were also at increased risk for persistent substance dependence, but compared with adolescents with 4+ risks, their struggles with dependence were more likely to be time-limited (Supplementary Fig. S2). The screening itself could serve as the basis for a brief intervention, as the cumulative risk index provides a powerful means of communicating to adolescents their overall risk in clear and understandable terms. Adolescents with 2 risks were seven times more likely than their peers with 0 risks to struggle with persistent dependence through early midlife. Adolescents with 3 risks were nine times more likely than their peers with 0 risks to have persistent dependence. For some adolescents, information about their risk status might motivate behavior change towards a lower risk category and prevent initiation or escalation of substance use. For adolescents who have 2 or 3 risk factors, and at least one of those risk factors includes frequent substance use, feedback about risk status might be combined with brief motivational interviewing and/or harmreduction strategies focused on substance use. For adolescents with symptoms of depression or conduct problems, brief interventions targeting these symptoms might be appropriate (e.g. behavioral activation for depression or brief behavioral parenting strategies for disruptive behavior).

Our results may well prove useful in designing and evaluating intensive prevention/early intervention programs in the future. Researchers planning these programs need to know approximately what proportion of their treatment group would develop persistent dependence. This information could be used to calculate the 'number needed to treat' to prevent one case a measure that is increasingly used to gauge the effectiveness of an intervention. For example, if adolescents with 4+ risks were the target of intervention, 60% would otherwise develop persistent substance dependence in adulthood (Table 4, PPV). Given a perfectly effective intervention, the 'number needed to treat' to prevent one case would be 1.67 (1/.60 = 1.67)(Cook & Sackett, 1995). As the proportion of the treatment sample who would otherwise develop persistent dependence decreases, the number needed to treat increases. Thus, prevention/early intervention studies that cast too wide a net in defining their treatment group have, from the start, limited their treatment's effectiveness.

The results of this study should be interpreted in the context of its limitations. First, although we selected some of the best predictors of substance dependence, prediction might be marginally improved by adding childhood and adolescent risks. Our sensitivity analyses showed that, with exception of adolescent frequent tobacco use, adding or subtracting a predictor did not make a difference in accuracy of the cumulative index. Moreover, each additional predictor yields diminishing returns while lengthening assessment (Ware, 2006). Substituting different predictors from the ones we examined here could lead to improvements in prediction, but multiple datasets will be needed to fully explore this possibility to avoid overfitting the model to this cohort. Substantial improvement in prediction may be unlikely however, because prediction is already quite good. A more promising strategy for improving upon prediction might be to include more predictors in models that could reveal combinations of risk factors that predict an especially high risk of persistent substance dependence (e.g. decision tree, cluster analysis, or neural network models).

A second limitation is that our findings are based on a single New Zealand cohort and require replication in independent samples. We view this report as proof of concept for using a cumulative risk index as an actuarial tool. The next steps include testing and refining the cumulative risk index in contemporary cohorts and in special populations, such as Native American adolescents, as well as testing the accuracy of the cumulative risk index in predicting persistent substance-use disorder, as defined by DSM-5.

In summary, we developed a universal screen for persistent substance dependence that (1) is based on population-representative data, (2) estimates total risk briefly and inexpensively by incorporating a relatively small number of well-established risk factors, and (3) aggregates risk factors using a simple algorithm. Although findings are preliminary, they suggest that we can predict with considerable accuracy which adolescents in the general population will struggle with persistent substance dependence in adulthood. The cumulative risk index may be simpler to use, less costly, and more accurate (AUC = 0.80) than more comprehensive screens, and these features are important considerations in universal screening. For example, an extensive profile of neuropsychosocial risk, including measures of brain and cognitive function, predicted adolescent binge drinking with an AUC of 0.75 (Whelan et al. 2014). We also showed that the cumulative risk index compared favorably to current risk assessment approaches, which narrow in on substance use as the primary risk indicator. Moreover, an adapted version of the cumulative risk model for use in community settings (a version that simply summed readily obtained adolescent risks) yielded an AUC of 0.79. Additional research is needed to validate the cumulative risk index, evaluate its practical utility, and address potential ethical issues that may be raised by screening adolescents for persistent substance dependence (Carter & Hall, 2011; Hall et al. 2015). The results presented here represent a first step toward establishing population-representative estimates of risk for persistent substance dependence that may be useful in research and practice.

Supplementary material

For supplementary material accompanying this paper visit http://dx.doi.org/10.1017/S0033291715002482.

Acknowledgements

We thank the Dunedin Study members, their families, Unit research staff, and Study founder Phil Silva. The Dunedin Multidisciplinary Health and Development Research Unit is supported by the New Zealand Health Research Council and New Zealand Ministry of Business, Innovation, and Employment (MBIE). This research received support from US-National Institute on Aging (AG032282, AG048895); UK Medical Research Council (MR/K00381X); and ESRC (ES/M010309/1). Additional support was provided by the Jacobs Foundation. Wayne Hall was funded by a National Health and Medical Research Council Australia Fellowship. Daniel W. Belsky was supported by the National Institute on Aging (T32-AG000029).

Declaration of Interest

None.

References

- **APA** (1987). *Diagnostic and Statistical Manual of Mental Disorders*, 3rd edn, revised. American Psychiatric Association: Washington, DC.
- **APA** (1994). *Diagnostic and Statistical Manual of Mental Disorders*, 4th edn. American Psychiatric Association: Washington, DC.
- Brook JS, Lee JY, Brown EN, Finch SJ, Brook DW (2011). Developmental trajectories of marijuana use from adolescence to adulthood: personality and social role outcomes. *Psychological Reports* **108**, 339–357.
- Carter A, Hall W (2011). Addiction Neuroethics: The Promises and Perils of Neuroscience Research on Addiction. Cambridge University Press: Cambridge.
- Chassin L, Flora DB, King KM (2004). Trajectories of alcohol and drug use and dependence from adolescence to adulthood: the effects of familial alcoholism and personality. *Journal of Abnormal Psychology* **113**, 483–498.
- Chung T, Smith GT, Donovan JE, Windle M, Faden VB, Chen CM, Martin CS (2012). Drinking frequency as a brief screen for adolescent alcohol problems. *Pediatrics* 129, 205–212.
- Clark DB, Chung T, Martin C (2006). Alcohol use frequency as a screen for alcohol use disorders in adolescents. *International Journal of Adolescent Medicine and Health* **18**, 181–188.
- Cook RJ, Sackett DL (1995). The number needed to treat: a clinically useful measure of treatment effect. *British Medical Journal* **310**, 452–454.
- Copeland WE, Angold A, Shanahan L, Dreyfuss J, Dlamini I, Costello EJ (2012). Predicting persistent alcohol problems: a prospective analysis from the Great Smoky Mountain Study. *Psychological Medicine* **42**, 1925–1935.
- **Costello A, Edelbrock C, Kalas R, Kessler M, Klaric S** (1982). *Diagnostic Interview Schedule for Children (DISC)*. National Institute of Mental Health: Rockville, MD.
- D'Agostino RB, Vasan RS, Pencina MJ, Wolf PA, Cobain M, Massaro JM, Kannel WB (2008). General cardiovascular risk profile for use in primary care - the Framingham Heart Study. *Circulation* 117, 743–753.
- Elley W, Irving J (1976). Revised socio-economic index for New Zealand. New Zealand Journal of Educational Studies 11, 25–36.

Evans GW, Li D, Whipple SS (2013). Cumulative risk and child development. *Psychological Bulletin* **139**, 1342–1396.

Felitti VJ, Anda RF, Nordenberg D, Williamson DF, Spitz AM, Edwards V, Koss MP, Marks JS (1998). Relationship of childhood abuse and household dysfunction to many of the leading causes of death in adults - the Adverse Childhood Experiences (ACE) Study. *American Journal of Preventive Medicine* 14, 245–258.

Fergusson DM, Horwood LJ, Ridder EM (2007). Conduct and attentional problems in childhood and adolescence and later substance use, abuse and dependence: results of a 25-year longitudinal study. *Drug and Alcohol Dependence* 88, S14–S26.

Grant BF, Dawson DA (1998). Age of onset of drug use and its association with DSM-IV drug abuse and dependence: results from the National Longitudinal Alcohol Epidemiologic Survey. *Journal of Substance Abuse* **10**, 163–173.

Grant BF, Goldstein RB, Saha TD, Chou SP, Jung J, Zhang H, Pickering RP, Ruan WJ, Smith SM, Huang B (2015). Epidemiology of DSM-5 alcohol use disorder: results from the National Epidemiologic Survey on Alcohol and Related Conditions III. JAMA Psychiatry 72, 757–766.

Grant BF, Hasin DS, Chou SP, Stinson FS, Dawson DA (2004). Nicotine dependence and psychiatric disorders in the United States - Results from the National Epidemiologic Survey on Alcohol and Related Conditions. *Archives of General Psychiatry* 61, 1107–1115.

Hall WD, Carter A, Yücel M (2015). Ethical issues in the neuroprediction of addiction risk and treatment response. In *Handbook of Neuroethics*, (ed. J. Clausen and N. Levy), pp. 1025–1044. Dordrecht, The Netherlands: Springer.

Heyman GM (2013). Quitting drugs: quantitative and qualitative features. Annual Review of Clinical Psychology 9, 29–59.

Hussong AM, Jones DJ, Stein GL, Baucom DH, Boeding S (2011). An internalizing pathway to alcohol use and disorder. *Psychology of Addictive Behaviors* **25**, 390–404.

Kendler KS, Jacobson KC, Prescott CA, Neale MC (2003). Specificity of genetic and environmental risk factors for use and abuse/dependence of cannabis, cocaine, hallucinogens, sedatives, stimulants, and opiates in male twins. *American Journal of Psychiatry* **160**, 687–695.

Kendler KS, Myers J, Damaj MI, Chen XG (2013). Early smoking onset and risk for subsequent nicotine dependence: a monozygotic co-twin control study. *American Journal of Psychiatry* **170**, 408–413.

Kirisci L, Reynolds M, Carver D, Tarter R (2013). Quick screen to detect current substance use disorder in adolescents and the likelihood of future disorder. *Drug and Alcohol Dependence* **128**, 116–122.

Krueger RF, Hicks BM, Patrick CJ, Carlson SR, Iacono WG, McGue M (2002). Etiologic connections among substance dependence, antisocial behavior, and personality: modeling the externalizing spectrum. *Journal of Abnormal Psychology* 111, 411–424.

Levy S, Weiss R, Sherritt L, Ziemnik R, Spalding A, Van Hook S, Shrier LA (2014). An electronic screen for triaging adolescent substance use by risk levels. *JAMA Pediatrics* **168**, 822–828.

McGue M, Iacono WG, Krueger R (2006). The association of early adolescent problem behavior and adult psychopathology: a multivariate behavioral genetic perspective. *Behavior Genetics* **36**, 591–602.

Meier MH, Caspi A, Houts R, Slutske WS, Harrington H, Jackson KM, Belsky DW, Poulton R, Moffitt TE (2013). Prospective developmental subtypes of alcohol dependence from age 18 to 32 years: implications for nosology, etiology, and intervention. *Development and Psychopathology* 25, 785– 800.

Milne BJ, Caspi A, Crump R, Poulton R, Rutter M, Sears MR, Moffitt TE (2009a). The validity of the family history screen for assessing family history of mental disorders. *American Journal of Medical Genetics, Part B: Neuropsychiatric Genetics* **150B**, 41–49.

Milne BJ, Caspi A, Harrington H, Poulton R, Rutter M, Moffitt TE (2009b). Predictive value of family history on severity of illness: the case for depression, anxiety, alcohol dependence, and drug dependence. *Archives of General Psychiatry* **66**, 738–747.

Moffitt T, Caspi A, Rutter M, Silva P (2001). Sex differences in Antisocial Behaviour: Conduct Disorder, Delinquency, and Violence in the Dunedin Longitudinal Study. Cambridge University Press: Cambridge, UK.

Moffitt TE, Caspi A, Taylor A, Kokaua J, Milne BJ, Polanczyk G, Poulton R (2010). How common are common mental disorders? Evidence that lifetime prevalence rates are doubled by prospective versus retrospective ascertainment. *Psychological Medicine* **40**, 899–909.

Odgers CL, Caspi A, Nagin DS, Piquero AR, Slutske WS, Milne BJ, Dickson N, Poulton R, Moffitt TE (2008). Is it important to prevent early exposure to drugs and alcohol among adolescents? *Psychological Science* **19**, 1037–1044.

Odgers CL, Milne BJ, Caspi A, Crump R, Poulton R, Moffitt TE (2007). Predicting prognosis for the conduct-problem boy: can family history help? *Journal of the American Academy of Child and Adolescent Psychiatry* **46**, 1240–1249.

Pardini D, White HR, Stouthamer-Loeber M (2007). Early adolescent psychopathology as a predictor of alcohol use disorders by young adulthood. *Drug and Alcohol Dependence* 88, S38–S49.

Pepe MS, Janes H, Longton G, Leisenring W, Newcomb P (2004). Limitations of the odds ratio in gauging the performance of a diagnostic, prognostic, or screening marker. *American Journal of Epidemiology* **159**, 882–890.

Poulton R, Hancox R, Milne B, Baxter J, Scott K, Wilson N (2006). The Dunedin Multidisciplinary Health and Development Study: are its findings consistent with the overall New Zealand population? *New Zealand Medical Journal* **119**, 45–55.

Poulton R, Moffitt TE, Silva PA (2015). The Dunedin Multidisciplinary Health and Development Study: overview of the first 40 years, with an eye to the future. *Soc Psychiatry Psychiatr Epidemiol* **50**, 679–693.

Rice ME, Harris GT (2005). Comparing effect sizes in follow-up studies: ROC area, Cohen's d, and r. *Law and Human Behavior* **29**, 615–620.

Robins L, Cottler L, Bucholz K, Compton W (1995). Diagnostic Interview Schedule for DSM-IV. Washington University School of Medicine: St. Louis.

Robins LN, Helzer JE, Croughan J, Ratcliff KS (1981). National Institute of Mental Health Diagnostic Interview Schedule - its history, characteristics, and validity. *Archives* of General Psychiatry 38, 381–389.

Rutter M (1981). Stress, coping and development - some issues and some questions. *Journal of Child Psychology and Psychiatry and Allied Disciplines* **22**, 323–356.

Sameroff AJ, Seifer R, Barocas R, Zax M, Greenspan S (1987). Intelligence quotient scores of 4-year-old children social-environmental risk-factors. *Pediatrics* 79, 343–350.

Stone AL, Becker LG, Huber AM, Catalano RF (2012). Review of risk and protective factors of substance use and problem use in emerging adulthood. *Addictive Behaviors* 37, 747–775.

Subramaniam GA, Volkow ND (2014). Substance misuse among adolescents: to screen or not to screen? *Jama Pediatrics* 168, 798–799.

Vanyukov MM, Kirisci L, Moss L, Tarter RE, Reynolds MD, Maher BS, Kirillova GP, Ridenour T, Clark DB (2009). Measurement of the risk for substance use disorders: phenotypic and genetic analysis of an index of common liability (vol 39, pg 233, 2009). *Behavior Genetics* **39**, 596–596. Ware JH (2006). Statistics and medicine - the limitations of risk factors as prognostic tools. *New England Journal of Medicine* 355, 2615–2617.

Weissman MM, Wickramaratne P, Adams P, Wolk S, Verdeli H, Olfson M (2000). Brief screening for family psychiatric history - the family history screen. Archives of General Psychiatry 57, 675–682.

Whelan R, Watts R, Orr CA, Althoff RR, Artiges E, Banaschewski T, Barker GJ, Bokde AL, Buchel C, Carvalho FM, Conrod PJ, Flor H, Fauth-Buhler M, Frouin V, Gallinat J, Gan G, Gowland P, Heinz A, Ittermann B, Lawrence C, Mann K, Martinot JL, Nees F, Ortiz N, Paillere-Martinot ML, Paus T, Pausova Z, Rietschel M, Robbins TW, Smolka MN, Strohle A, Schumann G, Garavan H, the IC (2014). Neuropsychosocial profiles of current and future adolescent alcohol misusers. Nature 512, 185–189.

Wilson PWF, D'Agostino RB, Levy D, Belanger AM, Silbershatz H, Kannel WB (1998). Prediction of coronary heart disease using risk factor categories. *Circulation* 97, 1837–1847.

Wright BRE, Caspi A, Moffitt TE, Miech RA, Silva PA (1999). Reconsidering the relationship between SES and delinquency: causation but not correlation. *Criminology* 37, 175–194.