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Research Article

## Do socioeconomic mortality differences decrease with rising age?

## Rasmus Hoffmann

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# Do socioeconomic mortality differences decrease with rising age? 

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#### Abstract

The impact of SES on mortality is an established finding in mortality research. I examine, whether this impact decreases with age. Most research finds evidence for this decrease but it is unknown whether the decline is due to mortality selection. My data come from the US-Health and Retirement Study (HRS) and includes 9376 persons aged 59+, which are followed over 8 years. The variables allow a time varying measurement of SES, health and behavior. Event-history-analysis is applied to analyze mortality differentials. My results show that socioeconomic mortality differences are stable across ages whereas they decline clearly with decreasing health. The first finding that health rather than age is the equalizer combined with the second finding of unequally distributed health leads to the conclusion that in old age, the impact of SES is transferred to health and is stable across ages.


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## 1. Introduction

The impact of the socioeconomic status (SES) on health and mortality has been observed many times in mortality research. Poor groups of people have generally two to three times higher death rates than rich ones (Auerbach and Krimgold 2001:31). The difference in life expectancy for Dutch men between the highest and lowest educational group is 4 years (Stronks 1997:3). In the 1980s, white men in the USA with a family income lower than $\$ 10.000$ had a 6.6 years lower life expectancy than those with an income higher than $\$ 25.000$ (Smith 1999:147). Within-country differences are at times much higher than international differences, e.g. the male mortality rate of those aged under 65 is higher in Harlem, New York, than it is in Bangladesh (McCord and Freeman 1990).

In spite of overall decreasing mortality levels, economic growth and improvements in medicine, mortality differentials between income groups and educational groups increased at ages 25 to 64 between 1960 and 1986 in the USA (Pappas et al. 1993:103). In the 1980s, this was also the case for men in all countries for which data are available, e.g. the United States, the Nordic countries and France (Valkonen 2001:8826).

Increasing differences here means increasing relative differences. Absolute differences, by contrast, may have decreased because of the overall declining level of mortality. However, results for the USA indicate that lower class mortality did not decline at all, which means that even absolute differences have increased (Auerbach and Krimgold 2001).

There is an ongoing debate about the causality of these mortality differentials. Some authors assume that health inequalities arise prior to differences in SES and that especially in later working ages the health status translates into SES via the ability to work (Smith 1999). Smith (2003 and 2004) finds that in pre-retirement ages new health events have a strong negative impact on income and wealth. In the following, I will leave this question aside and assume, based on the majority of research findings, that the main direction of causality especially in old ages goes from SES to health and that the notion of "SES impact on health and mortality" is thus generally justified (Fox et al. 1985, Goldman 2001). The concrete pathway of this impact will be further illustrated below.

While socioeconomic differences in health and mortality are well established by research findings, it is unclear whether these differences are stable across the life course or whether they decrease in old age. The latter is the most common finding made by researchers and has been explained using the following arguments:

1. Aging works as a leveler of social differences because biological processes assume dominance over social determinants and eventually everybody must die, regardless of social class (Liang et al. 2002:295).
2. The welfare state reduces socioeconomic differences in old age through benefits and social policy.
3. The impact of past experiences that are responsible for health differences, e.g. working conditions, fades out at old age.
4. The observed mortality differences get smaller in old age on the aggregate level because the surviving population is more homogeneous due to selective mortality.

The last argument means that the impact of SES on mortality on the individual level can be stable or even increases with age. This opposite result is based on the following assumptions:
5. The impact of past unhealthy experiences, e.g. unhealthy working conditions and smoking, is postponed to older ages.
6. Past experiences, e.g. education, accumulate and may interact with other factors, e.g. economic and social capital. The health outcome of this accumulation is incorporated into the "health stock".
7. Vulnerability increases in old age and makes differential exposures more harmful (House et al. 1994:221).

For a discussion of arguments 2 to 6 , see Ross and Wu (1996:107). For a more detailed elaboration of the cumulative advantage/disadvantage theory, see Crystal and Shea (1990), O'Rand (1996) and Dannefer (2003). As stated above, most research finds decreasing socioeconomic mortality differences with rising age (see e.g. Liang et al. 2002 and Marmot and Shipley 1996). But only few researchers have addressed this question in detail, some having attempted to identify a possible bias because of mortality selection and other problems of the measurement of mortality differences across different ages. House et al. (1994) find generally converging mortality differences but, depending on the indicator for SES, also an inconsistent pattern in the age trajectory of mortality differences. They stress the possibility that mortality selection causes convergence. After applying a method to account for a sample selection bias, Beckett (2000) concludes that the observed convergence is not biased by mortality selection. Ferraro and Farmer (1996), by contrast, show that mortality selection biases the measurement of gender and racial health differences over age. Ross and Wu (1996) reveal diverging health differences with increasing age; Fox et al. (1985) show similar social mortality differences below and above age 75. Otterblad

Olausson (1991) finds divergence for women and convergence for men in Sweden, which is similar to the recent results by Huisman et al. (2004), who show in an international comparison that relative mortality differences by education and housing tenure did not decline with age for women in some countries.

The aim of this paper is to gain deeper insight into the age pattern of socioeconomic mortality differences, to find possible reasons for such an interaction with age and also for the inconsistent previous research results on this topic. The theoretical background of this research is the question whether the interplay between social and biological factors in determining health and mortality of an individual is changing over the life course or not. Another related question is how we can understand social inequality in old age. Health may become so important for the living conditions and quality of life that it becomes an important aspect of social inequality.

To motivate the choice of variables and models in the empirical part of this paper, I will give an overview of possible causal pathways from SES to mortality. According to the majority of studies, material factors are responsible for a large part of socioeconomic mortality differences (e.g. Schrijvers et al. 1999). Money can buy healthy food, good housing, better medical treatment and other goods that are directly or indirectly relevant to maintaining a good health status. Education is important to receive knowledge about health risks and healthy behavior and in providing cognitive skills for dealing with complex information such as the association of behavior on one's personal health and the structures of the health care system. Social capital is helpful when a person needs information, connections and emotional and practical help.

Stress and behavior are factors that are on an intermediate level between SES and mortality. Stress is likely to be higher and health behavior is poorer in lower status groups (Stronks 1997, Auerbach and Krimgold 2001). Finally, on the societal level the health care system is an important factor that has an influence on whether a low status causes poor health and consequently higher mortality or not (Kunst 1997).

## 2. Data

My data come from the US Health and Retirement Study (HRS) and the study of Assets and Health Dynamics among the oldest old (AHEAD). These are two representative studies conducted by the Institute of Social Research (ISR), University of Michigan, and supported by the National Institute on Aging (NIA). They were started separately in 1992 and 1993 respectively and then combined in 1998, with a follow-up every second year (Soldo et al. 1997). Since HRS focuses on retirement ages
and AHEAD on ages of 70+, I merged them with the help of some data sets prepared by RAND (for information, see http://hrsonline.isr.umich.edu).

This resulted in a sample of 9376 persons born before 1934 (aged 59 to 107) surveyed from 1992 to 2000, with 2608 deaths during observation. I excluded black persons from the analysis because the small number of them in the sample would only show general racial mortality differences, which is not the purpose of this study. By the same token, it would not be possible to analyze their specific age trajectory of social mortality differences.

Institutionalized persons were already excluded in the original baseline sample but surveyed in the institution during the follow-up interviews. This may cause a bias. For example, single persons, persons with poor health and women are more likely to be in a nursing home and thus they are more likely to be underrepresented in the sample (Grundy and Sloggett 2003:936). Huisman et al. (2003) tested this bias and found that samples that exclude institutionalized persons underestimate socioeconomic health differences in older ages. The HRS sample, however, only omits them at baseline but follows them in the institutions. Thus, from wave to wave the percentage of people living in a nursing home comes closer to the percentage in the US population so that in the HRS wave of the year 2000 the differences are negligible. It is unlikely that my results are biased substantially by this slight under-representation.

The original HRS sampling procedure over-samples the population of Florida. I do not use special weights to compensate for this because there is no reason to believe that the Floridian population is systematically different from the US population beyond the characteristics that the numerous variables in the models control for. When the research focus is on multivariate modeling with relative risks as the outcome and not on descriptive statistics for the finite population, weights are shown to have a small impact on the results (Hoem 1989).

Missing values were almost entirely imputed when the data were prepared and combined by RAND. Rules for and results of this imputation can be found on the homepage cited above. When my data comes directly from HRS datasets and information for single waves are missing, I imputed it in a straightforward manner using information from the previous wave, or, if applicable, the mean of two waves.

## 3. Variables

The variables allow a detailed and time varying measurement of SES, health status and some control variables. Except for education, parents' mean age at death and having children all variables in the following list are time varying.

Education is measured in years of education (levels: 0-7,8-15,16+).
Wealth includes all assets of the household in which the person lives (bank account, real estate, shareholdings etc.) and is measured on three levels: lowest quartile, second lowest quartile and above median wealth.

Income is the net annual household income divided by a weighted number of persons living in the household (net equivalent income). The weight is 1 for the first person and 0.7 for all other persons in the household. Income is measured on the same three levels as wealth: lowest quartile, second lowest quartile and above median income.

Parents' mean age at death is the mean age at death of both parents (levels: -75, $76+$ ). Under certain conditions, it captures the genetic constitution that is transferred from parents to their children; see discussion section.

Children is an indicator for any own children (levels: yes, no). This variable measures partly social capital, i.e. whether a child is likely to look after the old person. However, it can not just be treated as a social status variable as it measures many different things. For example, having numerous children is an indicator of low social status and may be a cause for higher mortality whereas having no children may be a consequence of bad health (Doblhammer 2000).

Labor force status. This variable differentiates between working, being retired/disabled and not being in the labor force. While the labor force status is to a large extent a function of age and health (which I control for by using other variables), it additionally captures information on social status and every-day life - information that is predicting mortality.

Marital status is not a social status measure in a strict sense but it is related to SES. Firstly, marital status depends partly on social status, e.g. persons with a low social status are more likely to live alone (Goldman et al. 1995, O'Rand 1996). Secondly, marital status has a high impact on the social status in the sense that divorce or widowhood is often followed by a loss of economic and/or social capital. Moreover, marital status has an influence on health and mortality independent of SES. In this analysis, I combine the divorced with the never married persons because these are both very small groups that show a similar level of mortality.

Health behavior is an additive index focusing on three items that have shown to be important correlates of health: 1. physical activity (the persons were asked if they
engage in strong physical activity once a week or more), 2. ex-smoker, 3. current smoker. From the resulting four different categories of this score ( -1 to 2 ), the last two (with the worst health behavior) have been collapsed into one category because both of them were small.

Self-rated health. The question on self-rated health is posed with the traditional five categories: excellent, very good, good, fair, poor. I merged the first two categories.

Objective health is another additive index that includes four items: 1. in a hospital for more than 10 days per year, 2. limitations in activities of daily living (ADL), 3. a body mass index (BMI) at baseline $<21.4$ for men and $<19.5$ for women (=lowest decile), 4. loss of weight of more than 10 per cent of the body weight between two waves (= two years). From the resulting five different categories of this score (0 to 4), the last two (with the worst objective health) have been collapsed into one category because both of them were very small.

In principle, these items are reliable and objective descriptors of the health status, but the information as such is based on the respondents answer and not on objective measurements or tests. For more information about ADL, BMI and mortality, see Reuben et al. 1992, Losonczy et al. 1995, Himes 2000 and Greenberg 2001.

It was necessary to construct indices for health behavior and objective health because, given the limited number of cases, all interesting variables for the dimension of health would be too numerous to be included into the model.

Age is controlled for using four age groups (59-69, 70-79, 80-89, 90+)
Some variables have been tested in previous models and then skipped because they did not show significant results after controlling for other variables. The omitted variables are: occupational group, parents' education, going to church, children living nearby, drinking, high BMI, a gain in body weight of 10 per cent and more. I also checked and found that period or cohort effects do not bias the presented results.

To illustrate the frequencies for all categories of the variables, Table 1 shows the number of persons-years separately for men and women. In event-history-analysis, this shows information that is more exact than that revealed by the number of cases. Table 2 shows the corresponding persons-years for the different age groups used in the analysis.

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Table 1: Persons-years for the categories of the variables

|  |  | male | female |
| :---: | :---: | :---: | :---: |
| parents' age at death | -75 | 11645 | 15564 |
|  | 76+ | 11876 | 13751 |
| education | 0-7 | 2456 | 2688 |
|  | 8-15 | 16312 | 23249 |
|  | 16+ | 4752 | 3377 |
| children | no | 1872 | 3073 |
|  | yes | 21649 | 26241 |
| labor force status | work | 6442 | 3895 |
|  | retired/disabled | 16954 | 18591 |
|  | not in labforce | 124 | 6829 |
| marital status | married | 19263 | 13953 |
|  | widowed | 2846 | 12830 |
|  | divorced/never | 1413 | 2532 |
| wealth (percentiles) | 0-25 | 4600 | 8721 |
|  | 25-50 | 5452 | 7337 |
|  | 50-100 | 13469 | 13256 |
| income (percentiles) | 0-25 | 8339 | 14480 |
|  | 25-50 | 5940 | 6645 |
|  | 50-100 | 9242 | 8189 |
| health behavior | good | 2466 | 4987 |
| (act,exsmoke,smoke) | fair | 9356 | 15558 |
|  | poor | 11699 | 8770 |
| self rated health | excel/very good | 8694 | 10638 |
|  | good | 7536 | 8854 |
|  | fair | 4823 | 6372 |
|  | poor | 2468 | 3450 |
| objective health | excel/very good | 16663 | 18603 |
| (hospital,adl,thin,loss) | good | 5095 | 7713 |
|  | fair | 1450 | 2502 |
|  | poor | 313 | 497 |
| sum over all categories of one variable: |  | 23521 | 29315 |

## Table 2: Persons-years in the age groups

| Age | male | female |
| :--- | ---: | ---: |
| $59-69$ | 9612 | 8044 |
| $70-79$ | 8668 | 11775 |
| $80-89$ | 4618 | 7934 |
| $90+$ | 623 | 1562 |
| sum: | 23521 | 29315 |

## 4. Methods

I apply event-history-analysis with a model for the force of mortality as the outcome variable. The models include a baseline for the basic time variable age that is piecewise linear and coefficients for the multiplicative impact of categorical variables on the baseline risk of dying. The risks are computed as rates, based on occurrences (deaths) and exposures (persons-months) for specific combinations of variable levels. The results are displayed as rate ratios. I used STATA 8 and aML 2.04. The baseline for age covers the age range from 59 to 107 whereas the observation period is only 8 years, namely from 1992 to 2000. Thus, the cohorts are not real cohorts but partly synthetic ones in the sense that in spite of the longitudinal data, no individual in the data set is really observed from age 59 to ages above 67 .

The analysis of selective mortality is limited by the fact that only persons who survived until age 59 are included in the study. Persons who entered the study after age 59 are left-truncated, i.e. we only consider the period at risk after the respondents have entered the sample. STATA allows taking into account left-truncated cases by distinguishing between "time under risk" and "time under observation".

Different models are used in different steps to draw conclusions about the causal relationships between the predictor variables and their impact on mortality. Relative mortality rates are computed using different interactions and a term for unobserved heterogeneity. The general formula for the model is:

$$
\ln \mu_{i}(x)=y(x)+\sum_{j} \alpha_{j} X_{i j}+\sum_{k} \beta_{k} W_{i k}(x)+\sum_{\{l, m:(l, m) \neq(1,1)\}} \chi_{l m} I\{A=l, B(x)=m\}+U_{i}
$$

where $y(x)$ is the baseline hazard that depends on age, $X$ is a matrix of time constant variables and W denotes a matrix of time-varying variables that depend on age. The fourth term represents an interaction between a time constant and time varying
variable, A and B , where $I$ is an indicator that equals 1 for a specific combination of the levels of the two variables and that equals 0 otherwise. $U$ stands for a heterogeneity term that is assumed to measure an individually constant frailty in the sample. The mean and the variance of U in the population are assumed to decrease with age because mortality selection makes the population more homogeneous.

## 5. Results

Table 1 shows the relative risks of dying. The underlying models are without interactions and separate for men and women. The absolute baseline risk for the four age groups is not shown. The baseline risk roughly doubles from one age group to the next, i.e. with every ten years of age. The mean age at death is 79 for males and 84 for females.

Model 1 only contains the univariate results of each variable separately. All variables show the expected association with mortality and all of them are significant, except marital status for women and having children for men. Surprisingly, men with 8 to 15 years of education do not have a significantly lower mortality compared to those with 0 to 7 years of education.

In Model 2, all variables that directly or indirectly describe SES are included simultaneously while health variables are excluded. Naturally, the mortality differences between the levels of most of the variables get smaller than in Model 1 but, e.g. income and wealth still have an independent significant impact on mortality. We see that when wealth and income are controlled for, higher education no longer has a positive separate impact. Men with an intermediate level of education even have a significantly higher mortality than lowly educated men (see discussion below). Having children reduces mortality for women but not for men. Further, the retired, the disabled and persons who are not in the labor force have a higher mortality than those who still work.

Table 3: Event history model for mortality

|  |  | MODEL 1 |  | MODEL 2 |  | MODEL 3 |  |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  |  | male | female | male | female | male | female |
| parent's age at death | -75 | 1 | 1 |  |  | 1 | 1 |
|  | 76+ | 0.86*** | $0.77^{* * *}$ |  |  | 0.92 | 0.87** |
| education | 0-7 | 1 | 1 | 1 | 1 | 1 | 1 |
|  | 8-15 | 0.94 | $0.78{ }^{* * *}$ | 1.20** | 0.92 | $1.37^{* * *}$ | 1.03 |
|  | 16+ | 0.59*** | 0.63(***) | 0.99 | 0.86 | $\left.1.31{ }^{\star *}\right)$ | 0.94 |
| children | no | 1 | 1 | 1 | 1 | 1 | 1 |
|  | yes | 0.93 | 0.83** | 0.98 | 0.85** | 0.99 | 0.87* |
| labor force status | work | 1 | 1 | 1 | 1 | 1 | 1 |
|  | retired/disabled | $2.48^{* * *}$ | $3.36 * * *$ | $2.24 * * *$ | $3.02{ }^{* * *}$ | $1.54 * * *$ | $2.17^{* * *}$ |
|  | not in labor force | $3.17 * * *$ | 1.83 *** | $2.54 * *$ | $1.63^{* *}$ | $1.97 * *$ | 1.20 |
| marital status | married | 1 | 1 | 1 | 1 | 1 | 1 |
|  | widowed | 1.05 | 1.10 | 0.95 | 0.90 | 1.01 | 0.91 |
|  | divorced/ never | 1.46 *** | 1.17 | 1.25** | 0.80* | 1.22* | 0.77** |
| wealth (percentiles) | 0-25 | 1 | 1 | 1 | 1 | 1 | 1 |
|  | 25-50 | 0.88* | $0.71^{* * *}$ | 0.92 | $0.78{ }^{* * *}$ | 1.05 | 0.91 |
|  | 50-100 | $0.54 * * *$ | $0.57(* * *)$ | $0.65 * * *$ | $0.72{ }^{(* * *)}$ | 0.87 (*) | 0.90 |
| income (percentiles) | 0-25 | 1 | 1 | 1 | 1 | 1 | 1 |
|  | 25-50 | 0.75 *** | 0.60*** | 0.86** | $0.67^{* * *}$ | 0.95 | $0.75{ }^{* * *}$ |
|  | 50-100 | $0.52^{* * *}$ | $0.54\left(^{(* *)}\right.$ | $\left.0.722^{(* *}\right)$ | $0.65{ }^{(* * *)}$ | 0.82(**) | $0.74{ }^{(* *)}$ |
| health behaviour | good | 1 | 1 |  |  | 1 | 1 |
| (act, exsmoke, smoke) | fair | $2.21^{* * *}$ | $3.34 * * *$ |  |  | $1.73{ }^{* * *}$ | 2.40 *** |
| self rated health | excel/ very good | 1 | 1 |  |  | 1 | 1 |
|  | good | 1.58*** | 1.65*** |  |  | $1.32^{* * *}$ | $1.44 * *$ |
|  | fair | 2.60 *** | $2.68{ }^{* * *}$ |  |  | $1.85{ }^{* * *)}$ | 1.92(***) |
|  | poor | $6.11^{* * *}$ | $4.52^{* * *}$ |  |  | $3.38 * * *$ | $2.6{ }^{* * *)}$ |
| objective health <br> (Hospital,adl, thin, loss) | excel/ very good | 1 | 1 |  |  | 1 | 1 |
|  | good | $2.08 * * *$ | $1.76{ }^{* * *}$ |  |  | $1.36{ }^{* * *}$ | $1.22{ }^{* * *}$ |
|  | fair | 3.56 (***) | 3.43 *** |  |  | 1.74 (***) | 1.98*** |
|  | poor | $5.03\left({ }^{* * *)}\right.$ | 4.77(***) |  |  | 2.27 (***) | $\left.2.39{ }^{* * *}\right)$ |

* $: p<0.1 ; \quad{ }^{* *}: p<0.05 ; \quad$ *** $: p<0.01$

Stars in brackets mean that the parameter value is significantly different from 1 but not from the previous variable level.
Model 1 contains the univariate results of each variable separately,
Model 2 is multivariate including indicators for SES,
Model 3 adds the health variables and parents' age at death to Model 2.

Widows do not display a significantly different mortality from married persons. Men who are divorced or who have never married have a higher mortality whereas women in the same group have a lower one. Interestingly, the relative mortality risk of divorced or never married women turned from an insignificantly higher mortality according to the univariate results of Model 1 to a significantly lower mortality risk in Model 2 (see discussion).

Finally, income and wealth both have a strong diminishing impact on mortality.
One intermediate step between Model 2 and 3 is not shown here: it adds only health behavior to the SES variables and shows that the measured items of health behavior (physical activity, being an ex-smoker and being a smoker) changes the coefficients only slightly and do not remove the significance of any socioeconomic variables. This means that socioeconomic mortality differences to a large extent can not be explained by physical activity or smoking.

Model 3 is the full model, where the three health variables and also parents' mean age at death are added. We see that a high parents' mean age at death significantly reduces the mortality of women, and this supports the assumption that common genes in a family contribute to longevity. This interpretation is likely to be true, not least because the inclusion of parents' education in the model as an indicator of their social status does not change the impact of their age at death (results not shown). Thus, in Model 3 parents' SES is not a common background factor influencing both the parents' mean age at death and the mortality of the respondent. The genetic explanation is less likely if one assumes that parental education is not a good indicator for childhood conditions or if intergenerational continuities like smoking are more important than genes.

In the full model, wealth is no longer significant but most of the other socioeconomic mortality predictors still are. This indicates that the transition from a given health status to death is also influenced by SES. In the modeling of complex processes like those between SES, health and mortality, it is likely that some of the variables are intermediate variables for others and that they are not independent from each other. In this study, I find the highest correlations between wealth and income ( $\mathrm{r}=$ 0.47 ) and between wealth and education ( $\mathrm{r}=0.40$ ). As to the health variables, there are very low correlations between objective health and health behavior (men: 0.09 , women 0.16 ), low correlations between health behavior and self-rated health ( 0.22 and 0.14 ) and strong correlations between self-rated health and objective health ( 0.39 and 0.44 ). For SES as for health, it is clear that we have to measure multiple interrelated dimensions. This is justified as long as the different variables reveal interesting differences in their model results and these results are interpreted with caution.

The results in Table 1 are based on the assumption that the relative risks below or above 1 reduce or increase mortality as a multiplicative factor over the whole age
range and across all levels of different other variables. This assumption is too simplistic. Therefore, the following interactions give a more accurate picture of the influence of selected variables.

Figure 1: Female mortality with interaction between education and wealth (based on Model 2, low education and low wealth $=1$ )


An interaction between education and wealth (Figure 1) shows that for women material wealth is only beneficial when combined with middle or higher education and higher education is beneficial only in combination with at least average wealth. When I use income instead of wealth in such a graph, the result is similar. This means that beyond the result of Model 2, where the financial variables removed the positive influence of higher education, we now see that for women these two different resources have a complementary impact on mortality, i.e. both are necessary to have a mortality advantage.

In Figure 1, only the data points below the 0.8 line are significantly lower than 1 on the 5 per cent level and the decrease of the red line (for persons with 16 or more years of education) is significant. The mortality difference between persons with the highest and lowest education is only significant in the wealthiest group on the 10 per cent level. This shows that the overall level of significance is comparatively low due to an insufficient sample size. But the pattern of significance nevertheless supports the above interpretation, which claims that significant mortality differences occur among
wealthier persons and also when we compare wealth groups within the higher educated group.

The corresponding graph for males is not shown and discussed here. This is because it shows a different pattern, which is dominated by the surprising excess mortality of men with intermediate education. Since I do not know the reason for this mortality pattern (see discussion), an interaction between education and wealth for men would not provide deeper insights.

To address my central question, i.e. whether socioeconomic mortality differences are stable or declining with increasing age, it is necessary to run interactions between age, i.e. the basic time variable of the model, and a variable for SES. In the following analysis, I will use income as an indicator for SES. This is because it has the highest independent impact on mortality (see Table 1). In order to limit the material in this article, the following figures show mortality patterns for men. Women show neither more nor less convergence with age, but generally have an even less clear pattern over age than men.

Figure 2 shows the mortality for men with interaction between age and income. Note that the graph does not show the increase of mortality with age but only the relative differences between the three income groups.

Figure 2: Male mortality with interaction between age and income (based on Model 2, low income = 1)

age

As we saw in Table 1, men with the highest income have a significantly lower mortality. The upper bound of the confidence interval for the richest group (red line) for the four age groups is $0.84,0.99,0.95$ and 1.16 . The confidence interval for the oldest group is wider because of low case numbers in this group. Those with a middle income also display a lower mortality, but this is not significant at the 5 per cent level. Far from significant in this graph are the fluctuations of differences over age groups. This suggests that mortality differences between income groups are relatively stable over age and obviously not declining with increasing age. Again, the level of significance is not satisfactory, but here the differences in the oldest age group are non-significant because of the wide confidence interval due to low case numbers rather than because of a mortality convergence in old age.

Figure 3 repeats Figure 2 (thin lines) and shows the same interaction but based on Model 3, which controls for the health variables (thick lines).

Figure 3: Male mortality with interaction between age and income (based on Model 3, health controlled (HC), low income = 1)


We see that when we control for health, the lines for middle and higher income get closer to the reference line. This effect is limited to younger age groups, with the consequence that mortality differences between poor and middle/high income groups tend to increase with age. But due to the small sample, this increase is still far from being significant.

The next step is to address the problem of heterogeneity. Unfortunately, the model represented by the equation above, which includes the heterogeneity term U , did not show the expected results. Neither aML nor STATA 8 was able to identify heterogeneity in the estimation procedure. This is most likely due to the sample size, an insufficient observation time or insufficient variation in time varying variables and not to the absence of heterogeneity in the sample because even in models with very few variables heterogeneity was not found.

The assumption is that after controlling for heterogeneity, the relative risks would show a more realistic picture of determinants for individual mortality. The mortality differences between social groups as shown in Figure 2 and 3 are likely to be underestimated in higher ages because the population at high ages is more homogenous due to selective mortality. Thus, the true change of the impact of income over age for the individual can only be shown after a successful estimation of heterogeneity that in this case has to be postponed until better data are available. But since it is known in what direction the heterogeneity bias works, it is possible to conclude from the results so far that the impact of SES is not decreasing with rising age.

The next step shows how a health decline is affecting the impact of social status on mortality. Figure 4 shows an interaction between self-rated health and income. Age is still controlled for with four age groups, as it is in all models.

Figure 4: Male mortality with interaction between income and health (based on Model 3, low income = 1)


This interaction shows that income matters a lot when the person is in good health and that it has no impact when the person is in poor health. This means that poor health levels socioeconomic mortality differences. The mortality difference between the lowest and the other income groups is only significant at the 5 per cent level when people are in good health (RR: 0.45 , CI: $0.32-0.62$; RR: 0.69 , CI: $0.47-$ 0.95).

If the health status is so important for the impact of SES on mortality, then the resulting question is whether the health decline with age is equally distributed between social groups, enough to result in a leveling of the mortality between social groups. I would like to report here three aspects of health distribution. First, health declines generally with age: The correlation between age and average health during the study is $0.20^{* * *}$ for self-rated health and $0.34^{* * *}$ for objective health. There seems to be an adjustment for age in the self-estimation of health, which results in a lower correlation with age compared to the objective measure. But despite of the general health decrease with increasing age, health is unequally distributed between income groups: Table 2 shows the other two aspects of the health distribution: first, the average self-rated health status at the beginning of the observation and, second, the experience of a health deterioration, both by the three income groups from above. A transition from good to bad health here means that at the beginning of the observation period a person was in either the best or the second best category of either self-rated or objective health and has moved down at least two levels by the end of observation.

Table 2: Distribution and deterioration of health in different income groups by age

|  | age in 1992 | $\mathrm{n}=$ | low <br> income | middle <br> income | high <br> income |
| :--- | :--- | :--- | :--- | :--- | :--- |
| Percentage having very good <br> or good health at the beginning <br> of observation | $59-68$ | 3140 | 58.2 | 78.4 | 88.7 |
|  | $79-102$ | 4114 | 54.9 | 74.9 | 80.6 |
|  | $59-68$ | 2122 | 52.6 | 69.8 | 73.3 |
| Percentage that experiences a <br> health deterioration | $69-78$ | 2799 | 13.1 | 9.6 | 6.5 |
|  | $79-102$ | 1273 | 22.6 | 23.9 | 19.1 |

[^1]It is difficult to measure how large health differences are and even more so to measure how these differences change with age. But it is obvious that even if health generally declines with age, people with lower income initially have a lower health status and are more likely to experience a health decline. The number of cases for the analysis of health decline is smaller than that for the analysis of health at onset because only healthy persons can be considered for a possible health decline. In the oldest age group (the last row of the table), healthy persons are especially rare and selected, which may explain that the differences are not significant.

Concerning the question whether socioeconomic mortality differences decline with age or not, it is, finally, important to see whether the impact of the health status on mortality is stable across age groups. Figure 5 shows the interaction between age and health.

Figure 5: Male mortality with interaction between age and self-rated health (based on Model 3, very good health = 1)

age

The interaction between age and self-rated health reveals that mortality differences between health groups are very large in younger age groups (age 59 to 69). When all socioeconomic variables and the other health variables are controlled for, men with a poor self-rated health status at this stage have more than a six-fold higher mortality than those with very good health ( $\mathrm{RR}=6.6, \mathrm{CI}=4.4-9.8$ ). These mortality differences converge very strongly in older age groups. This convergence, and the convergence in Figure 4, is not due to self-estimation by the respondents because the
same interactions based on the objective health measure show an even stronger convergence (results not shown). Note that, naturally, it is possible to represent the interaction in Figure 5 in absolute terms. Mortality then would increase strongly with age and the distance between the lines, i.e. the absolute differences in the mortality risk, would only slightly decrease with age. But since I do not focus on the general increase in mortality with age, the chosen representation in Figure 5 is more appropriate.

The main result with respect to the convergence of socioeconomic mortality differences is that these differences are stable across age. However, they converge as health deteriorates. But health, in turn, assumes less importance for mortality in old age.

## 6. Discussion

### 6.1 Excess mortality of men with intermediate education

The surprisingly higher mortality for men with an intermediate education has been observed also elsewhere (e.g. Liang et al. 2002) and it has been interpreted as an educational mortality crossover due to selective mortality. An alternative explanation is that, holding income constant in the model, higher education means that the aforementioned education is not translated into higher income. This could be because the person never obtained a job that matches the educational level or he lost his job and thus experienced downward mobility, a move that may have been health related. This interpretation is supported by the fact that the excess mortality for middle educated men concentrates on the lower income and poorer health groups (results not shown). It also supports that education is not beneficial on its own but only when combined with higher income. This has been shown for women in Figure 1. One possible conclusion is that education as a measurement of SES has, besides several advantages, the disadvantage of being too stable across the life course.

### 6.2 Disadvantage for married women?

The finding that the relative mortality risk of divorced or never married women turned from an insignificantly higher mortality according to the univariate results of Model 1 to a significantly lower mortality risk in Model 2 may be due to an underrepresentation of institutionalized unhealthy women in the sample. But it is more likely to show a real disadvantage of married women. The scope of my presentation
does not allow for a detailed discussion of the reasons. But the fact that the sex difference emerges only after controlling for income and wealth may indicate that married women profit from higher material resources. Besides, they do not have an advantage or may even have a disadvantage when married net of the other factors in my analysis. Grundy and Slogett (2003) argued that women experience less of a disadvantage of being single than men because they engage less in unhealthy behavior in such situations (Johnson et al. 2000) and are more likely to substitute their singlehood with social networks (Goldman et al. 1995). In addition, they may even suffer in marriage, where they are likely to be the younger and healthier partner whose role it is to care for the ill spouse (Beckett et al. 2002).

### 6.3 Education and wealth work together to lower mortality

The interaction between education and wealth shows that for women neither education nor material wealth is beneficial separately (Figure 1). The excess mortality of women with high education but a low wealth status can be understood when we consider education as input and wealth as an output of the labor market. This group may suffer from not being successful in translating their education into material wealth or they lost their original occupational status. This would simply indicate that material wealth is a stronger mortality predictor than education. The other group with higher mortality, women with low education and a high wealth status, seems to indicate that this is not a general rule. Besides the order of importance between these dimensions of SES, there is a disadvantage of persons with an inconsistent social status, which means being on different levels in different dimensions of the social status (Siegrist et al. 1990). As indicated above, it was not possible to find or to reject this pattern for men because of the surprising higher mortality of men with an intermediate education, which makes the corresponding Figure for men difficult to compare to Figure 1.

### 6.4 SES mortality differences converge with declining health but not with age

The main result from the previous section, the convergence of socioeconomic mortality differences with worsening health but not with age, questions the majority of studies that do not separate age and health decline and find socioeconomic mortality differentials that converge with age.

The equalizing effect of worse health does not mean that social inequalities no longer exist after health has become poor. It rather raises the question to what extent health differences are caused by SES and if social inequality in old age is incorporated
in a more or less severe health decline and that, therefore, there is no longer social inequality in the transition from poor health to death. Thus, the question of social inequality in health is analogous to and becomes part of the question of social inequality in mortality.

Research findings reveal clear socioeconomic health differences at old age (e.g. Huisman et al. 2003) but the question of convergence or divergence with age is as unclear for health differences as it is for mortality differences. The majority of research results show converging health differences with age. Ross and Wu (1996) show that health differences increase up to age 90 and Huisman et al. (2003) more often find converging patterns but also mixed evidence depending on sex, country and indicator for SES. They studied socioeconomic inequalities in morbidity, using very good data from 11 European countries and find that morbidity differentials decrease with age in most countries but are stable and also increase in others. The convergence is less consistent for men than for women and less consistent for education as indicator for SES compared to income. The authors state that a small part of the observed convergence can be attributed to a bias caused by institutionalized persons, which have been excluded from their data. Concerning the obvious international differences, they assume that concrete risk factors, like drinking and smoking, may be responsible. Depending on the country, they may or may not cause a mortality convergence because such behavior is more common in lower status groups (which increases the gradient) and less common so among old people (resulting in a convergence with age).

In my study, I could only make an attempt to analyze health inequalities, which reveals increasing health differences because, from an already unequally distributed health at onset, the rate for health deterioration is also higher for low income groups (see Table 2 and Figure 6 below).

A closer look at Figure 3 shows that controlling for health decreases mortality differences in early old age. This is not surprising, since health is an intermediate variable and its inclusion in the model should, as observed in Model 3 in Table 1, decrease social mortality differences. The question arises why this inclusion does not change mortality differences in old age. A possible answer is provided in Figure 5: if the health status is less important for mortality in old age, controlling for health should accordingly change social mortality differences less so than in younger age groups.

My results allow a tentative evaluation of the arguments listed in the introduction, which leads to the following interpretation: Age increases for everyone, i.e. a convergence of socioeconomic mortality differences with age would actually indicate that the impact of SES decreases with age as a result of an equalizing welfare state policy or due to the temporal distance to unequal health experiences e.g. during work life (arguments 2 and 3 in the introduction). But instead, I find that socioeconomic mortality differences are stable across age groups (which supports arguments 5 to 7)
and that instead of age, poor health is the equalizer for social differences, maybe as a result of a universal shift from social to biological determinants of mortality when health decreases (argument 1). The arguments mentioned seem to be not mutually exclusive, e.g. accumulating social differences and the dominance of bad physical conditions over social conditions are possibly simultaneous.

The central question of my research, i.e. whether socioeconomic mortality differences decrease with age or not, has been answered by a modification of the question, namely the identification of two aspects of increasing age. Both of these aspects increase mortality but have very different implications for the impact of SES on mortality. The first aspect, increasing numerical age, seems to be trivial but, in fact, some of the arguments used to support the hypothesis of mortality convergence are based on numerical age. These arguments can now be rejected. The second aspect is declining health, where my finding that money matters less in poor health rejects the assumption that money is of major importance to people in bad health to get good treatment to prevent them from dying. Concerning declining health, the problem remains: the theoretically simple scenario that a socially mixed sample will experience a simultaneous health decline that would level social differences in mortality will practically never happen. The health decline of upper class persons will either be delayed, will start on a higher health level or will be slower. Therefore, it is difficult to say if the potentially leveling impact of a health decline is actually effective. This is because poor health is likely to be to a large extent the result of low SES and thus it is unequally distributed.

Figure 6 neglects the age dimension for a moment and shows the relation between health, SES and death, summarizing the findings from Table 2 (Transition A) and Figure 4 (transitions B and C).

## Figure 6: Transitions between good health and death



It is not obvious from my findings, how age is intervening in this constellation. On the one hand, Figures 2 and 3 show that the impact of social status is constant, on the other hand, the impact of health on mortality decreases with age (Figure 5). To answer this question, a very good measurement of socioeconomic health differences across age groups and maybe a multi-process model would be advantageous, but both go beyond the scope of this study at the present stage.

An advantage of my analysis is the measurement of social status and health on many dimensions. This is especially important in old age where, except for the fact that the occupational status is less important compared to younger ages, it is not exactly known which dimensions define SES. Another prerequisite for good modeling of individual health trajectories would be an observation time longer than 8 years, covering a larger part of age range 59 to $90+$. This lack of observation time may also be the reason for the failure to identify unobserved heterogeneity. This is because for such a procedure in a one level data set, one needs sufficient variation in time-varying variables.

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[^0]:    ${ }^{1}$ Max Planck Institute for Demographic Research, Rostock, Germany

[^1]:    Pearson's chi-square test has been applied to the original two-way tables (not shown) and the differences in the table are significant at the 99 per cent level except for the last row (see text).

