Antidiabetic Action of Bezafibrate in a Large Observational Database

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OBJECTIVE — The purpose of this study was to test the hypothesis that bezafibrate, an approved fibrate, can prevent or delay type 2 diabetes.

RESEARCH DESIGN AND METHODS — This was a retrospective cohort study using data from routine medical practice in the U.K., as captured by the General Practice Research Database (GPRD). Individuals chronically exposed to bezafibrate were compared with individuals chronically exposed to other fibrates. Hazard ratios (HRs) for incident type 2 diabetes were calculated using a Cox proportional hazards model. A post hoc analysis was used to examine the effect of bezafibrate on progression to use of oral antidiabetic medications or insulin in individuals with diabetes at baseline.

RESULTS — Bezafibrate users had a lower hazard for incident diabetes than users of other fibrates (HR 0.66 [95% CI 0.53–0.81]). This effect became stronger with increasing duration of therapy. Post hoc analysis of the effect of bezafibrate on progression of preexisting diabetes also showed a lower hazard for progression to use of antidiabetic medication (0.54 [0.38–0.76]) or progression to use of insulin (0.78 [0.55–1.10]).

CONCLUSIONS — Bezafibrate appears to have clinically important antidiabetic properties. Randomized controlled trials should be considered to assess the utility of bezafibrate in treating patients with diabetes or in preventing diabetes in high-risk patients.

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Type 2 diabetes is a major public health threat, expected to affect more than 221 million people worldwide by 2010 (1). One key target for diabetes drug development is the peroxisome proliferator–activated receptor (PPAR) (2,3). There are three isotypes that are of specific interest in metabolic diseases: PPAR-γ, PPAR-α, and PPAR-δ. The thiazolidinediones (e.g., pioglitazone) are PPAR-γ agonists used to treat diabetes through improvement of insulin response. The fibrates are PPAR-α agonists used to treat dyslipidemia by raising HDL and lowering triglycerides. PPAR-δ remains an investigational drug target with potential uses in diabetes, dyslipidemia, and obesity (4). Because dyslipidemia and diabetes are commonly comorbid, attempts have been made to create dual PPAR-α/γ agonists or pan-PPAR agonists, although none of these has reached the market (3).

In response to efforts to develop these agents, at least one observer has pointed out that the fibrate bezafibrate actually is a pan-PPAR agonist and affects insulin resistance (5). Post hoc analyses of a placebo-controlled randomized trial showed that bezafibrate may postpone or prevent type 2 diabetes (5,6). During a mean 6 years of follow-up, hazard ratios (HRs) versus placebo for incident diabetes were 0.59 (95% CI 0.39–0.91) in obese patients (5) and 0.70 (95% CI 0.49–0.99) in prediabetic patients (6). These clinical end point data were supported by biochemical evidence showing that bezafibrate slowed progression of insulin resistance (7).

Studies of the other fibrates (gemfibrozil, fenofibrate, clofibrate, and clofibrate) have not shown such effects (8–14), and these drugs are far more selective for PPAR-α than bezafibrate (15). Hence, it is reasonable to hypothesize that the status of bezafibrate as a pan-PPAR agonist may give it antidiabetic properties unique among fibrates.

Although not approved in the U.S., bezafibrate has been widely prescribed for dyslipidemia in the U.K. We used observational data to examine the a priori hypothesis that bezafibrate is unique among fibrates in reducing diabetes risk.

RESEARCH DESIGN AND METHODS — We conducted a cohort study using the General Practice Research Database (GPRD). Personal information was removed before inclusion in the database. The requirement for informed consent was waived by the University of Pennsylvania institutional review board and the GPRD Independent Scientific Advisory Committee.

Data source
The GPRD contains data abstracted from a computerized medical record system used by a subset of general practices in the U.K. Ninety-eight percent of the U.K. population receives all forms of health care through their general practitioners. The database is broadly representative of the U.K. population in terms of sex, age, and geography (16). We used data from 1988 through 2002.

The information prospectively collected in the database includes demographic information, all prescriptions written by the general practitioner, clinical diagnoses, specialty consultation notes, and hospital discharge diagnoses. Medical diagnoses are classified using Read Clinical Classification and the Oxford Medical Information System codes.

Participating general practices follow prospectively designed protocols for recording computerized clinical informa-
tion and uploading it to the research database. Data reaching predefined quality standards are so designated. More than 400 published epidemiological studies have been performed using the GPRD (16,17).

Study cohort
From all patients being followed in the GPRD, we included only person-time from individuals who were exposed to a fibrate. Individuals were only included who had been registered and up-to-standard with the GPRD for at least 12 months before initiation of the exposure drug, making this an inception cohort.

Because the primary outcome of interest was incident diabetes, any diagnostic code for diabetes or any use of home glucose-monitoring equipment or of drugs that are only used to treat diabetes (insulin, biguanides, sulfonylureas, thiazolidinediones, or acarbose) before the first fibrate prescription or within the first 90 days of fibrate therapy excluded that individual from participation in the primary analysis. The rationale for this exclusion was to avoid including prevalent diabetic subjects in the study cohort.

Exposure definition
The study group included those with more than one prescription for bezafibrate, as a way to identify those receiving chronic treatment. Because we excluded individuals developing diabetes within the first 90 days of therapy, we began follow-up with the 91st day of fibrate therapy. The duration of each prescription was either provided in the database or, when this information was missing, estimated from the number of pills dispensed. Exposure was assumed to continue 30 days after the end of the expected duration of the last prescription. Gaps over 60 days longer than expected between prescriptions were considered to mark a last prescription, although a patient could reenter the cohort with the next prescription. Clustering methods were used to account for single patients contributing multiple blocks of time to the cohort, and sensitivity analysis was done in which patients were censored at the first gap and not allowed to reenter the cohort.

Control groups were defined using parallel criteria, with person-time for control subjects defined by exposure to other fibrates without any history of bezafibrate use. The prespecified plan for this study was to maximize power by considering all nonbezafibrate fibrates as a single exposure and only distinguishing between individual fibrates in a secondary analysis.

Any patient who switched from one study group to another was censored at the time of the switch. In a secondary analysis, each specific fibrate constituted its own exposure group, compared with bezafibrate.

Outcome definition
The outcome of interest was clinical diagnosis of or treatment for diabetes, defined by at least two codes indicative of diabetes. Such codes included any diagnostic code for diabetes, any prescription for home glucose-monitoring equipment, or any prescription for insulin or an oral antidiabetic drug.

Post hoc analysis
In a post hoc analysis, two additional cohorts were created. These consisted of individuals who would have been eligible for the primary study but were excluded because of diabetes occurring before the 91st day of fibrate therapy. These individuals with baseline diabetes were divided into two groups. The new cohort consisted of individuals who had untreated diabetes at baseline (as identified by medical codes for diabetes or use of home glucose-monitoring equipment but no use of any antidiabetic medication). For this cohort, the outcome of interest was progression to use of any antidiabetic medication. In addition, individuals who were using oral antidiabetic therapy (a biguanide, sulfonylurea, thiazolidinedione, or acarbose) at baseline were treated as a separate cohort, with progression to use of insulin as the study outcome.

Statistical analysis
All exposure groups were first compared on baseline variables. For each exposure group, event rates were calculated. Next, Cox proportional hazard models were used to estimate unadjusted and adjusted HRs. Fully adjusted models were reported with all variables included in the model.

Covariates included the year that the exposure therapy was initiated. They also included a predefined list of factors known to be associated either positively or negatively with diabetes. These included sex, age, history of stroke, history of myocardial infarction, and use of the following drugs: ACE inhibitors, calcium channel blockers, β-blockers, thiazide diuretics, loop diuretics, and corticosteroids (18). These drugs were analyzed as baseline covariates and in sensitivity analysis as time-varying covariates. BMI and smoking status were available only for a portion of the population and were included only in secondary analyses. The presence of comorbidities was determined on the basis of identification of GPRD medical diagnostic codes in the year before the first fibrate prescription.

Five secondary analyses were performed: 1) comparison of bezafibrate users versus users of each individual fibrate; 2) stratification by duration of therapy; 3) stratification of bezafibrate users into approximate quartiles of average dosage (<200, 200–400, 400–600, and 600+ mg/day) with use of the low-dose category as a reference group; 4) restriction to subjects with baseline BMI data and incorporation of BMI into multivariable modeling; and 5) restriction to subject with baseline smoking data and incorporation of smoking into multivariable modeling.

RESULTS — Bezafibrate was used far more commonly (12,161 users) than any other fibrate (4,191 users). Of the other fibrate users, 1,465 used ciprofibrate, 502 used clofibrate, 824 used fenofibrate, and 1,400 used gemfibrozil. Baseline characteristics of bezafibrate users and other fibrate users were consistent with previously published research (Table 1) (18). Because of the large sample size, most baseline differences were statistically significant. However, few clinically significant differences were observed. Of interest, however, the prevalence of recorded obesity was very similar (5% vs. 6%) between the two groups. However, bezafibrate users were more likely to be female than other fibrate users (48% vs. 40%).

Users of all other fibrates were less likely to have baseline diabetes than users of bezafibrate (relative risk 0.90 [95% CI 0.82–0.98] adjusted for year of the first fibrate prescription). However, bezafibrate users were less likely to have diabetes before treatment initiation compared with only one subgroup, the fenofibrate users (1.25 [1.08–1.42] adjusted for year of first fibrate prescription).

Among bezafibrate users, 272 new cases of diabetes occurred, for an incidence rate of 8.5 cases per 1,000 patient-years (95% CI 7.5–9.5). Among users of the other fibrates, 131 new cases of diabetes occurred, for an incidence rate of 14.4 cases per 1,000 patient-years (12.1–17.1).
Table 1—Baseline characteristics and number of events in exposure groups

|                          | Bezafibrate | All other fibrates | P value for difference |
|--------------------------|-------------|--------------------|------------------------|
| n                        | 12,161      | 4,191              |                        |
| Person-years             | 32,091      | 9,067              |                        |
| Mean duration of use (years) | 2.6        | 2.2                | <0.0001                |
| Mode year of treatment initiation | 1993       | 1994               | <0.0001                |
| Age (years)              |             |                    |                        |
| 50                       | 20          | 22                 | 0.0112                 |
| 50–59                    | 33          | 33                 | 0.9840                 |
| 60–69                    | 37          | 33                 | <0.0001                |
| >69                      | 10          | 12                 | 0.0008                 |
| Male sex                 | 52          | 60                 | <0.0001                |
| History of myocardial infarction | 1           | 1                  | 0.7529                 |
| History of stroke        | 0           | 0                  | 0.9676                 |
| History of ACE inhibitor/angiotensin receptor blocker use | 5 | 6 | 0.0018 |
| History of calcium channel blocker use | 24 | 22 | 0.1756 |
| History of β-blocker use | 16          | 17                 | 0.2382                 |
| History of loop diuretic use | 5           | 5                  | 0.2178                 |
| History of thiazide diuretic use | 9          | 8                  | 0.2883                 |
| History of corticosteroid use | 3           | 3                  | 0.6388                 |
| Never smoker             | 19          | 21                 | 0.5114                 |
| Ever smoker              | 39          | 41                 | 0.5114                 |
| Not reported             | 42          | 38                 | <0.0001                |
| BMI                      |             |                    |                        |
| <25 kg/m²                | 9           | 8                  | 0.0449                 |
| 25–29.9 kg/m²            | 13          | 13                 | 0.6363                 |
| >29.9 kg/m²              | 5           | 6                  | 0.0035                 |
| Not reported             | 73          | 72                 | 0.0523                 |
| Number of cases of incident diabetes | 272       | 131                |                        |
| Cases/1,000 person-years (95% CI) | 8.5 (7.5–9.5) | 14.4 (12.1–17.1) | <0.0001 |

Data are % unless indicated otherwise. Individuals with baseline diabetes were excluded. History of cardiovascular events and drug use refer to history in the year before cohort entry. P values were generated using χ² and t tests.

Cox proportional hazard regression results are shown in Table 2. The unadjusted HR for the comparison between bezafibrate and all other fibrates was 0.58 (95% CI 0.47–0.72). Adjusting for year of treatment initiation attenuated the association slightly, yielding a HR of 0.64 (0.52–0.79). No other variables modified the point estimate by as much as 10%. The fully adjusted HR for incident type 2 diabetes was 0.66 (0.53–0.81, P = 0.0001). Analyses were repeated with stratification by year of treatment initiation, with no substantial change in the results or evidence of heterogeneity of results by year (data not shown).

Table 2—Prespecified secondary analyses consisting of HRs for exposure to bezafibrate

| Reference group          | HRs (95% CI) for incident type 2 diabetes in individuals exposed to bezafibrate | Unadjusted | Fully adjusted |
|--------------------------|---------------------------------------------------------------------------------|------------|---------------|
| All fibrate users        | 0.58 (0.47–0.72)                                                               | 0.66 (0.53–0.81) |
| Ciprofibrate users       | 0.53 (0.39–0.73)                                                               | 0.72 (0.52–0.99) |
| Clofibrate users         | 1.17 (0.63–2.14)                                                               | 0.78 (0.54–1.14) |
| Gemfibrozil users        | 0.30 (0.21–0.42)                                                               | 0.84 (0.46–1.55) |
| Fenofibrate users        | 0.81 (0.57–1.19)                                                               | 0.41 (0.29–0.58) |

Fully adjusted HRs are adjusted for year of treatment initiation, age, sex, history of congestive heart failure, history of myocardial infarction, and history of use of thiazide diuretics, loop diuretics, β-blockers, calcium-channel blockers, ACE inhibitors, angiotensin receptor blockers, or steroids.

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Table 3—HRs stratified by years of cumulative use

| Reference group | Year 1 (HR 95% CI) | Years 2–3 (HR 95% CI) | Years 4–5 (HR 95% CI) |
|-----------------|--------------------|----------------------|----------------------|
| All fibrate users | 0.71 (0.52–1.05) | 0.65 (0.45–0.90) | 0.51 (0.34–0.75) |

Fully adjusted HRs are adjusted for year of treatment initiation, age, sex, history of congestive heart failure, history of myocardial infarction, and history of use of thiazide diuretics, loop diuretics, β-blockers, calcium-channel blockers, ACE inhibitors, angiotensin receptor blockers, or steroids.

Table 4—Cox proportional hazard models for ad hoc analysis

| Reference group | For progression from unmedicated baseline diabetes to use of oral antidiabetic therapy | For progression from baseline use of oral antidiabetic therapy to use of insulin |
|-----------------|------------------------------------------|------------------------------------------|
| All fibrate users | 0.54 (0.38–0.76) | 0.78 (0.55–1.10) |
| Ciprofibrate users | 0.44 (0.28–0.69) | 0.78 (0.50–1.22) |
| Fenofibrate users | 0.57 (0.32–1.02) | 0.86 (0.52–1.77) |
| Gemfibrozil users | 0.74 (0.38–1.43) | 0.57 (0.31–0.95) |

All models treat bezafibrate as the exposure; reference group varies by row. Ciprofibrate was not used alone as a reference group because of an insufficient number of observations in the clofibrate group to support multivariable modeling. Fully adjusted models are adjusted for year of treatment initiation, age, sex, history of congestive heart failure, history of stroke, and history of drug use (ACE/angiotensin receptor blocker, calcium channel blocker, loop diuretic, thiazide diuretic, β-blocker, or steroid).

In vitro data and earlier post hoc analyses suggesting that bezafibrate can prevent or delay the onset of type 2 diabetes (5–7). It further indicates that this effect is unique to bezafibrate among the fibrates. These findings have important implications for research. Our findings are bolstered by the similarity of subjects in exposure groups on clinically relevant characteristics, the fact that the finding of a protective effect is of a clinically relevant magnitude, statistically significant, and robust to sensitivity analyses including adjustment for BMI, and the monotonic duration-response relationship.

The results of the post hoc analysis are reassuring. It was reasonable to worry that fenofibrate was more likely to be prescribed to individuals with a high risk for diabetes or unrecorded diabetes, creating a falsely elevated hazard for development of diabetes during fenofibrate treatment compared with bezafibrate treatment. It was hence useful to do a post hoc analysis confined to individuals who already had diabetes. In this post hoc analysis, bezafibrate also appeared to have antidiabetic properties.

Taken together, these findings support and complement previous observations. Post hoc analyses of the Bezafibrate Infarction Prevention (BIP) Study have suggested that bezafibrate may reduce the hazard for incident diabetes, with point estimates for the HR of 0.59 and 0.70 (3,6). The fully adjusted point estimate from our study (0.66, 95% CI 0.3–0.81) is very consistent with those earlier results. These additional results are important because they confirm a post hoc analysis in a new study with this as a prespecified hypothesis, they generalize the results to a broader population than the original post hoc analysis did, and they provide considerably more precise point estimates.

Another publication from the BIP showed that bezafibrate attenuated progression of the homeostasis model assessment of insulin resistance marker for insulin resistance in all patients, suggesting that bezafibrate might also slow progression of diabetes (7). Our results were consistent with that hypothesis as well, both for progression from diagnosis to any use of antidiabetic drugs (HR 0.54, 95% CI 0.38–0.76) and for progression from use of oral antidiabetics to insulin (0.78, 0.55–1.10). The finding for progression to insulin was a trend but was not statistically significant. In addition, the findings on diabetes progression should be noted to be post hoc and are not adjusted for multiple comparisons.

The major limitation of this study was the potential for unadjusted confounding, a problem in any observational study. Despite the similar indications for the fibrates, the drugs are clearly not identically prescribed. Most strikingly, bezafibrate was by far the most commonly used fibrate in the U.K. and had the highest proportion of female users. Of most concern, rates of baseline diabetes differed by exposure group. Adjusting for these baseline differences did not change our results. Further, it is reassuring that bezafibrate still appeared to be protective even when compared with ciprofibrate, clofibrate, or gemfibrozil (which were not preferentially prescribed to diabetic subjects compared with bezafibrate). It is especially reassuring that a post hoc analysis of diabetes progression that could not have been confounded by preferential prescribing to diabetic subjects still showed a protective effect from bezafibrate. We in turn note that this post hoc analysis was not part of the original study design, was subject to false-positive results due to multiple comparisons, and used rough proxies for diabetes progression (progression to oral or insulin therapy). No observational study can completely exclude the possibility of confounding by indication, and the results of this study need to be confirmed in a subsequent randomized study. Another potential limitation of this study is the likelihood that some incident cases of diabetes were not captured by the database; however, such misclassification would most likely be nondifferential and would bias any finding toward the null.

In summary, this study strongly supports the idea that bezafibrate can prevent type 2 diabetes, confirming a post hoc analysis in a prior study. The effect size estimates from this study are comparable to those reported for other studies assessing the use of thiazolidinediones and metformin to prevent diabetes (19). Given concerns about cardiovascular risk with existing oral antidiabetic agents (20,21), bezafibrate may offer a unique opportunity to treat or prevent diabetes while maintaining a favorable cardiovascular risk-benefit profile. However, it would not be appropriate to establish a new indication without randomized controlled trial data to confirm these findings, similar to those recently conducted to study the antidiabetic properties of coleselovam (22). In light of the increasing population risk of diabetes, a trial that could establish the effectiveness of an inexpensive and
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